**ABSTRACT**

**Introduction** Transdiagnostic cognitive–behavioural therapy (CBT) targets common psychological factors that underlie multiple disorders. While transdiagnostic interventions are a promising new approach, limited research has evaluated these treatments within the alcohol and other drug (AOD) sector for young people with comorbid mental health symptoms. This project will examine the feasibility and preliminary efficacy of FullFix—a new risk-targeted transdiagnostic CBT telehealth programme for comorbid AOD and depression/anxiety disorders in young people. Secondary aims are to identify moderators and mediators of treatment outcomes, to determine how and why treatment is effective and who is most likely to benefit.

**Methods/design** Participants will be 130 young people (aged 16–35) accessing AOD services in Queensland, Australia, with comorbid mental health symptoms. They will be randomised to receive either the FullFix intervention plus standard AOD care or standard AOD care alone. Primary outcomes on AOD use and mental health symptoms will be reassessed at 6 weeks, 3 months, 6 months and 12 months, along with secondary outcomes of emotion regulation, social connectedness, perceived self-efficacy, coping skills and quality of life. The trial commenced on October 2018 and expected completion date is September 2021.

**Ethics and dissemination** Ethical approval for this trial was provided by the University of Queensland (#2018001185). The results of the trial will be disseminated through publication in a peer-reviewed scientific journal, scientific presentations at conferences and distributed via a report and presentations to the partner organisation.

**Trial registration number** ACTRN12618001563257.

**INTRODUCTION**

Epidemiological and clinical studies consistently show that alcohol and other drug use (AOD) and mental disorders frequently co-occur.1–4 For instance, in a nationally representative survey of 8841 Australians (aged 16–85 years), one in three respondents with a past year substance use disorder also met criteria for an anxiety disorder, and one in five met criteria for an affective disorder.7 The high rate of co-occurring substance use and mental health disorders incur significant costs to society, and are a key challenge to policy and practice. Comorbidity is related to a more severe and chronic illness course, increased relapse risk, worse treatment outcomes, higher risk of suicide and greater social and functional impairment.8–11 This challenge is magnified in people accessing AOD services, who frequently present with additional psychosocial complexities, such
as history of trauma, neglect, homelessness, criminal justice involvement and unemployment.\textsuperscript{12,13}

High rates of comorbidity and associated treatment challenges are of particular concern in young adults. Mental health and substance use disorders are at their peak and the leading cause of disability and death worldwide in adolescence and young adulthood.\textsuperscript{14,15} Further, although there is strong epidemiological evidence that 75\% of AOD and mental health problems emerge before the age of 25, treatment typically does not occur till many years later,\textsuperscript{16,17} with one study finding an 18-year gap between problem identification and receipt of AOD treatment.\textsuperscript{18}

Current practice approaches often fail to address the complex patterns of comorbid mental health and AOD problems with which young people present. Although an integrated approach to the management of comorbidity is recommended,\textsuperscript{19–21} treatments that target comorbidity remain underutilised within AOD treatment services.\textsuperscript{22} FullFix, an integrated telehealth intervention for young people with comorbid AOD and mental health problems, was developed to address these treatment concerns.

Telehealth interventions
Few young people experiencing mental health problems, particularly AOD problems, access treatment. For instance, one study found fewer than one in four young people with substance use disorders had accessed health services in the past year.\textsuperscript{23} Young people face many barriers to accessing treatment, with stigma and shame being a key deterrent to help-seeking.\textsuperscript{24} Telephone-delivered (telehealth) interventions provide an innovative, youth friendly way of rapidly increasing treatment access to the 96\% of young people who own a mobile phone.\textsuperscript{25} Previous studies in adolescents reported young people found telephone-delivered interventions more acceptable, convenient, flexible and less stressful than attending a clinic.\textsuperscript{26,27} Further, meta-analyses have found that telephone-delivered psychological therapies, particularly cognitive–behavioural therapy (CBT), have promise in improving mental health outcomes.\textsuperscript{26–29} Within AOD treatment, telephone-delivered interventions have been shown to be effective for reducing risk of future AOD use and related harm in young people.\textsuperscript{30,31}

Transdiagnostic approaches
Transdiagnostic interventions, which cut across diagnostic boundaries to target risk factors underlying multiple problems, are a promising new approach to the treatment of comorbid disorders.\textsuperscript{32,33} Theoretically, transdiagnostic approaches suggest that diverse mental health disorders have common latent factors, which develop from interactions between common genetic, biological, psychological and environmental vulnerabilities.\textsuperscript{34–39} Common identified transdiagnostic risk factors and protective factors for both AOD include (1) impulsive, sensation seeking and neurotic personality traits,\textsuperscript{40–44} (2) coping, self-control and emotion regulation skills\textsuperscript{34,36,43,45–48} and (3) social factors, such as social support or the social normative environment.\textsuperscript{39,49,50} Developing an intervention that targets these three key areas could provide more efficient and effective treatments for AOD and comorbid mental disorders.

There is a growing evidence-base demonstrating transdiagnostic approaches, including transdiagnostic CBT,\textsuperscript{22,51–53} mindfulness-based interventions\textsuperscript{54} and emotion-regulation focused interventions\textsuperscript{45} have positive outcomes for depression and anxiety disorders. However, a recent narrative review of transdiagnostic approaches for comorbid depression and AOD disorders concluded that while transdiagnostic treatment approaches have been developed for emotional disorders and anxiety disorders, there are limited programmes targeting AOD comorbidities.\textsuperscript{22} The authors conclude that despite the lack of trials examining transdiagnostic interventions for AOD, such process-based approaches hold promise, as treatments that target transdiagnostic factors—specifically, negative affect, anhedonia, rumination, experiential avoidance, emotion regulation and distress tolerance—lead to improvements in depression and AOD use.

Objectives of the study
Overall, the evidence suggests that a transdiagnostic approach to addressing comorbid AOD and mental health problems has a strong theoretical basis with empirically derived treatment targets, and has the potential to improve clinical outcomes in clinically complex populations. Yet, there is a dearth of studies examining transdiagnostic interventions for young people with AOD use and comorbid problems. The current project aims to examine the feasibility and efficacy of a new risk and protective factor-targeted transdiagnostic CBT telehealth programme (FullFix) for comorbid AOD and mental health problems in young people delivered as an adjunct to standard AOD care, compared with standard AOD care alone. Feasibility and outcomes are assessed at 6 weeks (approximately mid-treatment), 5 months (post-treatment), 6 months and 12 months postbaseline.

Secondary aims are to identify moderators and mediators of treatment outcomes, to determine how and why treatment is effective and who is most likely to benefit. This information will be used to further refine FullFix to make it more targeted and effective.

We hypothesise that individuals receiving FullFix plus standard AOD treatment will achieve significantly greater improvements on the primary outcomes of AOD use and mental health as well as the secondary outcomes of mental well-being and functioning outcomes, compared with individuals receiving standard AOD treatment alone at 3-month, 6-month and 12-month postbaseline.

METHODS AND ANALYSIS
Study design
This multicentre randomised controlled trial (RCT) will compare the efficacy of a transdiagnostic CBT
intervention (FullFix) delivered as an adjunct to standard care, compared with standard care alone for young people accessing inpatient and outreach AOD services. The trial will also assess the feasibility to deliver the FullFix intervention in these settings. The trial is registered (ACTRN12618001563257) and follows Standard Protocol Items: Recommendations for Interventional Trials research protocol guidelines (see online supplemental materials) as well as Consolidated Standards of Reporting Trials guidelines (see figure 1).

**Study settings**

Young people (16–35 years) will be recruited from Lives Lived Well (LLW) AOD treatment services. This includes seven services providing outreach and on-site services and a residential inpatient treatment service in Queensland, Australia. Approximately 50% of clients referred to these services are presenting for treatment for the first time.

**Eligibility criteria**

Inclusion criteria are: (1) 16–35 years of age; (2) seeking help for current AOD use from participating LLW services; (3) Alcohol, Smoking and Substance Involvement Screening Test (ASSIST) score of either >10 for alcohol use, or >4 for illicit drug use; (4) a score ≥10 for either the Patient Health Questionnaire-9 (PHQ-9); or the Generalised Anxiety disorder (GAD-7); or a score of ≥3 on the Primary-Care Post-traumatic Stress Disorder Screen; and (5) access to mobile phone technology. Exclusion criteria are: (1) current serious medical

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**Figure 1** Projected CONSORT diagram, showing expected retention and loss. CONSORT, Consolidated Standards of Reporting Trials.
problem or traumatic injury; (2) not fluent in spoken or written English; (3) unmodified hearing impairment; (4) current acute high suicide risk (intention and plan) as assessed by the treating clinician; (5) current psychosis or diagnosed schizophrenia that is currently being treated with antipsychotics; (6) history of traumatic brain injury or organic brain disease that interferes with treatment delivery and (7) currently in acute alcohol or drug withdrawal. There are no restrictions on concomitant care or other interventions that participants can access or take part, in addition to the study trial.

Sample size calculations

Medium effects (f=0.25) are anticipated based on our previous CBT trial. With power set at 0.90, and alpha of 0.05, 52 participants in each group, or a total sample of 104, is required. We predict a 20%–30% attrition rate based on our previous work and will therefore aim to recruit a minimum of 132 (20% attrition) and up to 149 (30% attrition) participants. All participants who are randomised to a treatment condition will be included in the intent-to-treat analysis.

Recruitment and follow-up procedures

Participants will be recruited from new clients entering the service or beginning a new programme at a participating site. After enrolling in a treatment programme, LLW staff will provide potential participants (clients aged 16–35 seeking assistance for AOD use) with brief information about the research, and then obtain their verbal or written consent to be contacted by a researcher, as part of a standard consent process. Programme enrolments will be monitored weekly by a trained research assistant and the project manager to ensure the protocol is being followed. A research assistant will directly contact LLW staff to follow-up on referrals that do not have consent status (yes/no) recorded. Trained research assistants with a minimum honours degree in psychology will then contact the consenting client within 3 days to complete the screening measures and obtain full informed consent. Participants meeting inclusion criteria will be sent a survey link to complete the baseline assessment online. Those that complete the baseline survey will be randomised to receive either FullFix+standard AOD care or standard AOD care only.

Participants will be asked to complete follow-up surveys online at 6 weeks, 3 months, 6 months and 12 months post-baseline. All participants will be followed up for intent-to-treat purposes, regardless of treatment completion. Reminders to complete the survey will be sent to participants by research assistants blind to treatment allocation, and will be followed up over the phone if the survey is not completed within 1 week. A follow-up protocol has been developed, which includes a schedule for when to send reminders and via what modality (email, text messages to mobile devices, social media, phone-calls and locator phone-calls and text messages). Participants will be reimbursed AUD$20 for completing the baseline and each follow-up survey.

Recruitment commenced September 2018, with enrolment of the first participant occurring in October 2018. Recruitment was concluded in July 2020, with the last participant enrolment occurring in August 2020. Data collection is due to be finished in September 2021.

Randomisation

A randomisation sequence has been created using statistical software R with the package ‘blockrand’. The randomisation sequence was stratified by service area (six sites) and age (16–20, 21–25, 26–30, 31–35), with a 1:1 allocation using random block sizes of 4, 6 and 8 to randomise participants into (1) FullFix + standard AOD treatment or (2) standard AOD treatment alone. The randomised sequence is contained in a password protected spreadsheet. Randomisation sequence and allocation will be conducted by independent research assistants not involved in recruitment or data collection. Allocation will be communicated to the chief investigator and LLW staff to assist with treatment planning.

FullFix intervention

FullFix is an eight-session telehealth transdiagnostic CBT intervention which targets the risk and protective factors common for comorbid AOD and depression/anxiety problems. It incorporates the evidence based QuikFix personality-specific coping skills brief intervention. QuikFix is a two session intervention comprising assessment feedback/information, motivational interviewing (MI), psychoeducation and coping skills training targeting the participant’s predominant maladaptive personality traits related to problematic AOD use. It has been found to result in larger reductions in the number of standard alcohol drinks consumed at both 1 and 12 months follow-up among young people with alcohol related injuries and illnesses, compared with assessment feedback/information or MI alone. In the pilot RCT for QuikFix, although people who received QuikFix reported significantly reduced distress (on the Kessler Psychological Distress Scale (K10)), 88% of participants nevertheless continued to report at least moderate distress at 1-month follow-up (K10 score ≥17), suggesting that people with mental health comorbidity may benefit from a more intensive intervention. This is addressed in FullFix which is designed to be delivered using a stepped care framework. Specifically, all participants are first provided with the QuikFix brief intervention (sessions 1 and 2 of FullFix), followed by sessions focused on building protective factors and strengths (session 3) and teaching participants a self-regulation framework for implementing coping skills in different situations (session 4).

After these four core modules, young people are offered up to four additional, optional, sessions. The order of the four additional sessions is flexible and are determined in collaboration with the client. These modules target (1) positive and negative emotional
awareness and regulation, (2) healthy behaviours and behavioural regulation, (3) cognitive regulation and (4) social connectedness and support. These sessions are tailored to the individual’s specific risk and protective factor profile by a trained psychologist using a case formulation approach. All young people are offered the opportunity to receive these additional four sessions, regardless of their initial response to FullFix. Modules are delivered individually in 30–50 min telephone therapy calls for up to 8 weekly or fortnightly sessions, which may be broken into two shorter calls per week if the participant prefers. Table 1 provides a summary of all intervention modules.

An MI style (eg, empathically amplifying the person’s own concerns)65 is used throughout FullFix to: (1) build readiness and commitment to make a change (including pros and cons of current use and change); (2) negotiate change goals; and (3) develop a plan for change each session, with a focus on building self-efficacy to achieve change. Development of the plan includes developing specific, measurable, and achievable goals within a timeframe, discussing support and role-models for achieving goals, recalling past successes of behaviour change, personal strengths or strategies for achieving goals, potential barriers or challenges to implementing the plan (and ways to overcome barriers), and the reasons underlying the goals.

**Standard AOD care**

The control group receives standard AOD care in accordance with standard practice at the participating site. Details of episodes of care and session case notes will be recorded by LLW clinicians. Frequency of sessions, duration, modality (phone or face-to-face) and type of session (eg, counselling, case management) will be measured for each participant. An independent researcher will conduct a file audit to check the reliability of the episodes of care data for a random sample of 20% of participants to determine inter-rater reliability. Personalised assessment feedback on the participant’s AOD use and related physical, psychological and social consequences will also be sent to the usual care clinician via email.

**Community services**

Community standard AOD care typically consists of case management and AOD counselling, delivered via phone or face to face by an LLW AOD clinician (typically an allied health professional with at least a bachelor’s degree). A typical episode of care consists of 2–10 sessions. Usual care is not based on specific guidelines, nor does it have a prescribed number of sessions, frequency or duration of sessions; as this is determined on a case by case basis by the AOD worker. Typical treatment approaches used include CBT for substance use, MI, psychoeducation and counselling.

**Residential rehabilitation services**

Residential Rehabilitation standard AOD care typically consists of a 6-week treatment programme, with a maximum length of stay of 12 weeks, and telephone support by pretreatment/post-treatment AOD workers to assist with admission into the residential facility and the transition back to the community. The programme includes regular group sessions covering substance use (eg, triggers, cravings, building motivation to change, drug refusal skills), emotion regulation and management, grief and loss, developing a relapse prevention plan; one on one counselling/case management; as well as creativity, exercise, nutrition and relaxation activities.

**Patient and public involvement**

Consultations were conducted with clinicians and staff of the participating organisations to inform the study protocols and design. Patients were not involved in the design of the study or intervention. The project manager and relevant members of the research team meet regularly with team leaders and area managers of the services involved in the project, as well as monthly meetings with organisation management. At these meetings, feedback is invited on the conduct of the research. Adaptations to the protocol will be made, if necessary, based on this ongoing feedback. The measures assessing client feedback will be used and incorporated into future studies involving service users.

**Feasibility**

A number of outcomes will be used to assess feasibility. We will assess recruitment methods using the number of potential participants per week who are: eligible, approached, consented and who meet inclusion criteria. We will also assess the number of consenting participants who are willing to be randomised, the number who complete all baseline assessments within 14 days, and the number who complete follow-up assessments within 30 days of target date. Additionally, we will measure reasons for refusal or non-completion at all stages of research (screening, enrolment, randomisation and all follow-up time-points). To specifically assess feasibility of the intervention, we will record number of consenting participants who commence their randomly allocated treatment within 14 days, and sessions completed by participants.

**Measures**

The assessment schedule is presented in table 2.

**Demographics**

Participants will be asked to report their age, gender, occupation, years of education, relationship status, living arrangements (including postcode), ethnicity, family and personal history of mental and substance use disorders, history of traumatic brain injury or organic brain disease.
### Table 1  Overview of FullFix intervention

<table>
<thead>
<tr>
<th>Session</th>
<th>Content</th>
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| Preintervention assessment    | Assessment completed via phone or online. See table 2 for list of assessment measures, with measures relevant to the FullFix intervention marked with *.
| Core sessions                 |                                                                                                                                            |
| Module 1                      | Personalised assessment feedback on self-reported level of AOD use and how this compares to national norms, AOD-related problems, motives for AOD use and depression and anxiety symptoms. Patterns, goals and motives for AOD use are explored, across different contexts. Psychoeducation about harm minimisation strategies. Develop a harm minimisation plan in the form of an implementation intention, that is, 'if situation Y is encountered, I could cope more effectively by using Z, in order to achieve my AOD use goal'. |
| Module 2                      | Training in two cognitive behavioural coping skills targeting the participant's predominant impulsive personality style, measured by the S-UPPS-P: (1) negative urgency (tendency to act rashly under extreme negative emotions); (2) positive urgency (tendency to act rashly under extreme positive emotions); (3) sensation seeking (tendency to seek out novel and thrilling experiences); (4) lack of premeditation (tendency to act without thinking) — within the intervention termed impulsivity. Coping skills are orientated to whether the emotions have more of a depressive or anxious presentation. Mindfulness training in how to increase awareness of, observe and sit with emotional responses. Mindful breathing (if high in anxiety). Behavioural activation (if high in depression). Thought challenging. |
| Negative urgency              | Mindfulness training in how to increase awareness of, observe, and sit with emotional responses. Increase positive reinforcements not related to AOD. |
| Positive urgency              | Mindfulness training in how to increase awareness of, observe, and sit with emotional responses. Savouring techniques to increase awareness and ability to focus on and experience positive feelings from everyday activity (eg, mindful eating). Identify and schedule natural highs from functional activities (eg, exercise, skateboarding, rock climbing) and daily experiences (food, music). |
| Sensation seeking             | Mindfulness training to increase awareness of thoughts and emotions. Learn and apply the Stop-Think-Do rubric to help slow down decision-making processes sufficiently to consider likely outcomes of behaviours. |
| Lack of premeditation         | Assessment feedback on the strengths they reported using most frequently at baseline. Identify three strengths in themselves and when they have used strengths. Strength-spotting exercises to increase their ability to identify strengths in themselves and others: (1) think about an occasion when they were at their best, then describe what happened and what strengths they showed; or (2) to identify a person they admire and what strengths that person shows. |
| Module 3                      | Training in a self-regulation framework to decrease reactivity to events and increase regulatory behaviours, using a four step approach with the acronym GAGE: ► Get with it—Using the process of Breathe-Ground-Centre-Focus to bring mind to the present moment. ► Appraise—Appraise the situation and ask themselves what is going on and what can I do about it? ► Go!—Take goal-directed action, based on their appraisal of the situation. Act rather than react. ► Explore—Evaluate how their actions went by asking themselves what went well and what could be done differently next time. |
| Module 4                      |                                                                                                                                            |
| Additional sessions           |                                                                                                                                            |
|                               |                                                                                                                                            |
**Primary outcomes**

**Alcohol and drug use**
- The WHO ASSIST screener for problem/risky substance use, covering tobacco, alcohol, cannabis, amphetamine-type stimulants (including ecstasy), hallucinogens, opiates and 'other drugs'.

**Mental health**
- Depressive symptoms are assessed using the PHQ-9. This is a nine-item self-report measure that assesses depressive symptoms in the past two weeks and is suitable for use in both primary care settings and over the phone. Anxiety is assessed using the GAD-7, which has been validated across a number of populations and will be suitable for assessing anxiety in young adults.

**Secondary AOD outcomes**
- AOD-related problems: An adapted version of the 15-item Controlled Drinking Self-Efficacy Scale (CDSSE) will be used to measure current and task-oriented coping strategies. The CDSSE has been found to be reliable and valid in young adults.[6] AOD-related problems: Brief Young Adult Alcohol and Other Drugs Consequences Questionnaire adapted from Kahler and Strong.[68] will be used to assess AOD-related problems in different domains (e.g., social, medical, legal, family, vocational).

**Secondary well-being outcomes**
- Coping: The Emotion-Oriented and Task-Oriented subscales from the Coping Inventory for Stressful Situations scale will be used to measure emotion-focused and task-oriented coping strategies. Well-being: The brief 14-item version of the Mental Health Continuum-Short Form (MHC-SF),[70] will ask participants about their emotional, psychological and social well-being in the past month. The MHC-SF has been found to be reliable and valid in young adults.[31]

**Quality of life**
- The Brief Questionnaire of the Mental Health Continuum (MHSC-Q),[71] will ask participants about their emotional, psychological and social well-being in the past month. The MHSC-Q has been found to be reliable and valid in young adults.[32] The Brief Questionnaire of the Mental Health Continuum will collect information about the patient's quality of life (QOL) across a number of domains (e.g., social, medical, legal, family, vocational). The QOL assessment is then repeated three times across a number of populations and will be suitable for assessing anxiety in young adults.[60]

**Session Content**

**Social connectedness**
- Psychoeducation on the importance of social support and social connectedness. Exercise examining the social groups and connections participants have in their social network. Exercise exploring the type of relationship they have with each social connection, whether it supports or hinders their goals, and importance of each social connection. Explore how they would like their social network to look, and what support they would like to give and receive.

**Emotion regulation**
- Psychoeducation on emotions (what are emotions, function of emotions, thought-feeling-behaviour connection). Exercise on recognising and learning to observe and sit with emotional responses (positive and negative). Identify current maladaptive and adaptive emotion regulation coping strategies. Identify and practice emotion regulation strategies they would like to improve or use more frequently (expanding on the skills taught in session 2 and 4).

**Cognitive regulation**
- Psychoeducation on cognitions (thoughts vs facts, negative automatic thoughts, and thought-feeling-behaviour connection). Identify unhelpful thinking patterns and current unhelpful coping (e.g., suppression, rumination, and avoidance) and helpful coping (e.g., reappraisal, problem solving, and acceptance). Identify and practice strategies they would like to improve or use more frequently (expanding on the skills taught in session 2 and 4).

**Healthy behaviours**
- Awareness and understanding of behaviours that are important for health: specifically sleep, diet and nutrition, physical activity. Skills building in ways to regulate physiological responses, specifically through diaphragmatic breathing and relaxation. Practice specific coping skills for managing cravings (e.g., Urge Surfing), if experiencing urges or cravings.

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Table 1 Continued

<table>
<thead>
<tr>
<th>Session</th>
<th>Content</th>
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<tbody>
<tr>
<td>Social connectedness</td>
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<tr>
<td>Measures</td>
<td>Initial assessment</td>
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<td></td>
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<tr>
<td>Family history of mental health issues</td>
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<td>Relationship, employment and medical history</td>
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<td>Hearing impairment</td>
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<td>Alcohol withdrawal and previous AOD treatment in last month</td>
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<td>AOD use</td>
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<td>BQYADCOQ*</td>
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<td>GAD-7*</td>
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<td>EXITS*</td>
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<td>Signature strengths*</td>
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<td>FFMQ-SF</td>
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<td>Well-being and quality of life</td>
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<tr>
<td>MHC-SF</td>
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Continued
of domains. The EUROHIS-QOL has previously been used with Australian populations and found to be a reliable and valid measure.

### Transdiagnostic risk factors

#### Impulsivity

**Stressors**


- The 18-item Difficulties in Emotion Regulation Scale (Gratz KE, Roemer L. Emotion regulation, emotion dysregulation, and implications for psychopathology. J Psychopathol Behav Assess 2004;26(1):41–51). This study uses a modified version of this scale, the Drug Use Motive measure, to assess motives for AOD use.

- The 24-item Signature Strengths Survey will be used. This measures captures the degree to which a strength is seen as defining to the self and the self-rated top strengths (essential strengths) a person possesses.

#### Protective factors

- Social support will be measured using a four-domain, 24-item Signature Strengths Survey. The measure has good reliability and validity, and has been validated for use in young adults.

### Social factors

- Social support will be measured using a four-domain, 24-item Signature Strengths Survey.

### Other measures

- The 18-item Difficulties in Emotion Regulation Scale will be used to measure emotion regulation difficulties. The measure has high reliability and validity, and has been validated for use in young adults.

<table>
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<tr>
<th>Measures</th>
<th>Initial assessment</th>
<th>Follow-up assessments</th>
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<td></td>
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</table>

*Assessments are necessary for preintervention assessment for FullFix intervention.*

AOD, alcohol and other drug; ASI-3, Anxiety Sensitivity Index-3 Dimension; ASSIST, Alcohol, Smoking and Substance Involvement Screening Test; BOYAADCQ, Brief Young Adult Alcohol and Other Drugs Consequences Questionnaire; CDSE, Controlled Drinking Self-Efficacy Scale; CDSI, Controlled Drinking and Drug-taking Self-Efficacy; CISS, Coping Inventory for Stressful Situations; DERS-18, Difficulties in Emotion Regulation Scale-18 items; DUMM, drug use motivations measure; EXITS, Exeter Identity Transition Scale; FFMMOSF, Five-Facet Mindfulness Questionnaire-Short Form; GAD-7, Generalised Anxiety Disorder; MHC-SF, Mental Health Continuum-Short Form; OTI, Opiate Treatment Index; PEQ, Patient Experiences Questionnaire; PHQ-9, Patient Health Questionnaire-9; PTSD, post-traumatic stress disorder; QOL, European quality of life; SUPPS-P, Short Impulsive Behavior Scale.
Treatment satisfaction
To measure participants’ experiences with clinicians, the Patients Experience Questionnaire from the Improving Access to Psychological Therapies programme will be used at post-treatment assessment time-points. The scale is five items measuring client experiences on a 5-point Likert scale (1: strongly disagree to 5: strongly agree). Open ended responses will also be included to collect feedback from the participants about their level of satisfaction with their treatment and to better inform treatment procedures.

Treatment fidelity
Clinicians for the FullFix condition are registered psychologists on the research team, with a master’s degree in clinical psychology. Training and supervision for FullFix will be conducted throughout the trial by registered clinical psychology supervisors. To assess intervention adherence and fidelity, telephone treatment sessions will be audiorecorded, with the participant’s consent. Clinicians will also record a session component checklist for each session completed. Supervision meetings will be held fortnightly to monitor session delivery and treatment adherence. Additionally, the supervisor will review a randomly selected session audio segment, independently rating it and a Session Component Checklist, and discuss any departures from protocol. A random sample (20%) of session recordings will be independently rated for treatment fidelity to ensure core features of the allocated treatment are delivered.

Statistical analyses
Feasibility and acceptability outcomes will be assessed using descriptive statistics of rates of eligibility, enrolment, attrition, reasons for refusal and follow-up rates. Independent means t-tests or non-parametric equivalents (for non-normal data) and \( \chi^2 \) tests will be used to compare treatment completers and non-completers. Responses to open ended questions will be evaluated using content and thematic analysis, as appropriate based on the nature of responses, to identify themes and patterns across responses.

The independent variable is treatment group, with two levels: (1) FullFix +Standard AOD care and (2) Standard AOD care alone. The primary outcome variables are AOD use and related harm (ASSIST, OTI) and depression/anxiety symptoms (PHQ-9, GAD-7). Secondary outcomes include well-being (MHC-SF), quality of life (EURO-HIS-QOL), drug taking confidence and coping self-efficacy to resist using alcohol heavily (adapted CDSE). A series of mixed effects, repeated measures analyses of variance (with time as the within subjects factor and group as the between subjects factor) will be employed to determine whether there are treatment group differences in improvements in primary and secondary outcomes of young people at 6weeks, 3-month, 6-month and 12-month follow-up compared with baseline. Analyses will be conducted on an intention-to-treat basis with all participants included in analyses, regardless of whether they withdrew from the treatment programme or did not complete all follow-up assessments. Analyses will be adjusted for covariates (eg, demographic characteristics) and factors associated with missing data. Bootstrapping will be used to account for any violations in the assumption of normality of residuals.

Moderators of treatment effects and process outcome variables include transdiagnostic risk and protective factors (emotion regulation, impulsivity, social support, coping skills and self-perceived strengths). To identify mechanisms and moderators of change and the strength of the factors involved, both multilevel models and structural equation models (using analyses in \( R \) ) will be used for mediation and moderation analyses.

Ethics and dissemination
Ethical approval for this trial was provided by the University of Queensland (#2018001185). Safety and risk management protocols attend to safety or urgent treatment issues. A clinical trials committee meets every 5months to monitor the project’s implementation, clinical and research integrity. Urgent issues are determined via email or telephone consultations and recorded to ensure consistency. Any young person who reports the presence of significant suicidal ideation/intent is provided with additional telephone support by a research psychologist and referral to additional treatment (through participating research sites or external local services), and an afterhours crisis team (if required). Any safety or urgent treatment issues will be managed as per usual safety and risk management procedures by the LLW clinicians. If a client screens positive for suicide risk during the screening procedure, their caseworker is made aware, and they are followed up by LLW treating staff.

Dissemination
The results of the trial will be disseminated through publication in peer-reviewed scientific journal, scientific presentations at conferences and distributed via a report and presentations to the partner organisation.

CONCLUSION
The current paper reports the study protocol of a pilot RCT determining the feasibility and efficacy of a new telehealth transdiagnostic CBT intervention for young people aged up to 35 years, presenting to AOD treatment services.

Identifying treatments that can address comorbidities within substance use settings that are accessible and feasible is of high importance considering the (1) low rates of treatment among young-people, (2) high rates of comorbidity, (3) poorer outcomes and (4) social and economic burden of comorbidity. Results of this study are expected to increase access to evidence-based and cost-effective care for young people with comorbid AOD use and mental health problems.
Author affiliations
1 School of Psychology, The University of Queensland, Brisbane, Queensland, Australia
2 National Centre for Youth Substance Use Research, The University of Queensland, Brisbane, Queensland, Australia
3 School of Medicine and Public Health, The University of Newcastle, Callaghan, New South Wales, Australia
4 Rural Clinical School, The University of Queensland, Toowoomba, Queensland, Australia

Twitter Genevieve Dingle @Genevieve132

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ORCID iDs Zoe Walter http://orcid.org/0000-0001-8310-4021
Nina Pecuca http://orcid.org/0000-0003-0264-1680

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