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Research Paper

What Works in Youth Suicide Prevention? A Systematic Review and Meta-Analysis

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SUMMARY

Background: Young people require specific attention when it comes to suicide prevention, however efforts need to be based on robust evidence.

Methods: We conducted a systematic review and meta-analysis of all studies examining the impact of interventions that were specifically designed to reduce suicide-related behavior in young people.

Findings: Ninety-nine studies were identified, of which 52 were conducted in clinical settings, 31 in educational or workplace settings, and 15 in community settings. Around half were randomized controlled trials. Large scale interventions delivered in both clinical and educational settings appear to reduce self-harm and suicidal ideation post-intervention, and to a lesser extent at follow-up. In community settings, multi-faceted, place-based approaches seem to have an impact. Study quality was limited.

Interpretation: Overall whilst the number and range of studies is encouraging, gaps exist. Few studies were conducted in low-middle income countries or with demographic populations known to be at increased risk. Similarly, there was a lack of studies conducted in primary care, universities and workplaces. However, we identified that specific youth suicide-prevention interventions can reduce self-harm and suicidal ideation; these types of intervention need testing in high-quality studies.

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1. Introduction

Suicide is the second-leading cause of death among young people and rates appear to be increasing [1]. Suicidal thoughts and behaviors (defined as suicide attempt or self-harm with clear or unclear suicidal intent) are more common than suicide [2] and predict future suicide and suicide attempts [3], with the period following a first suicide attempt associated with highest risk [4]. Presenting to hospital with self-harm significantly predicts subsequent suicide in youth [5]; with the period immediately following discharge from psychiatric inpatient treatment associated with highest risk for suicide [6]. The period following hospital discharge therefore provides a crucial opportunity for intervention. Suicidal ideation is a necessary precursor to suicide attempt and as such also requires intervention. Although suicidal ideation is arguably a distinct concept from suicidal behavior, for ease of reading it is

included under the term “suicide-related behavior” throughout this review unless otherwise specified.

The majority of OECD countries have a national suicide prevention strategy and many identify young people as requiring specific attention [7–9]. In accordance with international best practice, most strategies recommend a comprehensive approach to suicide prevention spanning universal approaches (i.e., delivered to the whole population), selective approaches (i.e., delivered to groups or communities believed to be at higher risk of suicide) and indicated approaches (i.e., delivered to individuals displaying suicide-related behaviors). Strategies also recommend interventions operate across a range of settings, including clinical, educational, workplace and community settings [1]. More recently, strategies have called for interventions to be delivered in digital, as well as face-to-face, settings [10,11].

Strategies must encompass evidence-based interventions if they are to reduce suicide [1]. Generating such evidence in suicide prevention, however, is complex [12]. Statistically, suicide is a relatively rare event, therefore it is often unfeasible to obtain sample sizes necessary to demonstrate the impact of interventions on this outcome. Moreover,

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Research in Context

Evidence Before This Study

Prior to this study systematic reviews in suicide prevention have been limited by either only including RCTs, or by concentrating on particular settings (e.g., schools) or intervention type (e.g., gatekeeper training), and as such do not cover the full spectrum of approaches. The more comprehensive systematic reviews do not focus specifically on youth.

Added Value of This Study

This is the first systematic review and meta-analysis to synthesize the full spectrum of suicide prevention approaches in young people. It identified a large number of studies conducted across clinical, educational/workplace and community settings. Studies also tested the full spectrum of interventions including universal means restriction and educational interventions, selective interventions such as training programs, indicated interventions such as cognitive or dialectical behavior therapy, and multimodal interventions that combined education with either screening or gatekeeper training. The meta-analysis found that interventions delivered in both clinical and educational settings appear to have an impact on suicide-related outcomes at post-intervention and follow-up. In community settings, multifaceted, place-based approaches seem to have an impact on rates of suicide and self-harm. Overall, study quality was limited.

Implications of All the Available Evidence

The review identified that specific youth suicide-prevention interventions can reduce both self-harm and suicidal ideation in clinical, school and community settings, challenging the nihilism that often pervades in suicide prevention. Indeed, the number and range of studies identified by this review is encouraging and reflects increasing investment and best practice internationally when it comes to youth suicide prevention. However, there was an absence of studies conducted in low-middle income countries where large numbers of suicides occur, or with specific populations known to be at elevated risk of suicide, such as indigenous or same-sex attracted young people. Similarly, few studies were conducted in primary care, workplace or university settings, and very few utilized digital platforms. Additionally, many studies simply tested interventions that had previously been designed for adults as opposed to young people specifically. Together these findings suggest that important opportunities for youth suicide prevention are currently being missed. These gaps now need to be addressed by researchers, research funders, and by policy makers if we are to successfully address the rising rates of suicide among young people worldwide.

many interventions do not lend themselves to being tested using randomized controlled trials (RCTs), typically considered the gold-standard [13]. As such, researchers assess changes in other more prevalent outcomes, including self-harm and suicidal ideation, using alternative study designs. Therefore, when synthesizing the evidence regarding what works in youth suicide prevention, alternative study designs warrant consideration.

Whilst previous reviews have synthesized this evidence, many only include RCTs [14]. Additionally, many concentrate on particular settings

(e.g., schools) [15], or types of intervention (e.g., gatekeeper training programs) [16], and as such do not cover the full spectrum of approaches. Finally, systematic reviews that include a range of study designs and intervention types do not focus specifically on youth [17,18]. Hence, a comprehensive review of the literature on youth suicide prevention interventions spanning the range of settings, study designs and intervention types, is required to better understand what works in youth suicide prevention. This will help policy makers, clinicians, service providers and commissioners determine the focus of future suicide prevention efforts.

We conducted a systematic review and, where possible, meta-analysis, of all studies examining the impact of interventions that were specifically designed to reduce suicide-related behavior in young people. Overcoming the limitations of previous reviews, we placed no restriction on study setting, intervention approach, or study design.

2. Methods

The methodology was informed by the Cochrane Collaboration [19] and the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines [20].

2.1. Study Selection and Classification

2.1.1. Inclusion Criteria

Studies of any design were eligible for inclusion in this review, provided they: [1] evaluated the impact of an intervention specifically designed to reduce suicide-related behavior; [2] assessed a suicide-related outcome, including suicide, suicide attempt, self-harm (defined as intentional self-injury and/or self-poisoning where suicidal intent was either not specified or was unclear), suicidal ideation, suicide risk, and/or reasons for living; [3] targeted young people aged 12–25 and/or if data on young people (mean age between 12·0 and 25·0) was specifically reported; [4] were published in a peer-reviewed journal or identified via the reference lists of included articles; and [5] were written in English.

2.1.2. Exclusion Criteria

Studies were excluded from the review if: [1] they were not implemented with the expressed and primary purpose of preventing or reducing suicide-related behavior. Under this criterion, studies of indicated interventions were excluded if they did not recruit participants based on present or recent suicidal ideation or behavior. Additionally, studies of means restriction approaches were included only if the intervention was implemented, wholly or partially, to prevent suicide. As such, studies of firearm regulations implemented with the expressed and primary purpose of preventing homicide were excluded under this criterion. Studies were also excluded if they: [2] did not measure and report on a suicide-related outcome (as defined above); this included studies that exclusively measured *non-suicidal* self-injury, as this is generally considered to be a separate phenomenon; [3] did not target young people, or if data relating to outcomes for young people could not be disaggregated from that adults; [4] employed a non-experimental design; [5] were not published in a peer-reviewed journal; [6] were not available in English; or [7] did not contain any unique relevant data over and above the first included study.

2.2. Search Strategy

We searched Medline, PsycINFO, and EMBASE from January 1 1990 to September 21, 2017. Keywords relevant to suicide-related behavior, intervention type and youth were combined using standard Boolean operators (see Appendix). Key words were developed by consensus among the author group and in consultation with a librarian. In addition, we hand-searched the reference lists of all previous reviews retrieved via the search.

In the first instance study titles and abstracts were screened by five of the review authors (EB, JR, SH, NS, KW). Due to the large number of

studies retrieved two review authors independently screened 10% of the total number of records retrieved. Cohen's Kappa [21] was 0.748 and Prevalence-Adjusted and Bias-Adjusted Kappa (PABAK [22]) was 0.978, indicating excellent agreement regarding inclusion and exclusion of studies. Discrepancies were resolved by discussion. In the second stage of screening, full texts of potentially relevant studies were screened for inclusion by four authors (EB, JR, SH, NS). Full text double-screening was not undertaken, but review authors met regularly to resolve any queries.

2.3. Data Extraction and Classification

Data were extracted independently by seven authors (JR, EB, SH, NS, KW, DC, AM) using a pilot tested pro forma. The following information was extracted: (i) author(s) and publication date; (ii) country; (iii) study design; (iv) setting from which participants were recruited; (v) study sample or population characteristics; (vi) intervention description; (vii) details of control or comparison group (classified as treatment as usual (TAU), enhanced TAU and placebo), and; (viii) outcome data on suicide deaths, suicide attempt, suicidal ideation, suicide-related behavior, and/or self-harm at the point of post-intervention and (where appropriate) longest follow-up (note that follow-up periods varied). Where studies used more than one measure for an outcome, data from the measure that was most commonly used across all included studies were used, as has been done previously [23]. Two authors (SH and KW) undertook double data entry of all outcome data.

Studies were classified according to the following taxonomy. In the first instance studies were classified according to the setting from which the participants were recruited (i.e. clinical, education or workplace, and community). If participants were recruited from multiple settings, the study was classified according to the setting from which participants were primarily recruited. Studies were then classified by study design (i.e. RCTs and non-RCTs) and then by intervention approach (i.e. universal, selective, indicated). Some studies combined a number of different intervention approaches. In these cases studies were classified as 'multi-modal' when the intervention comprised a number of different components implemented together (e.g. psycho-education AND screening), and 'multiple' when studies tested the impact of different interventions that were implemented separately (e.g. psycho-education program in location A and gatekeeper training in location B). They were then classified according to intervention type (e.g. means restriction, educational, therapeutic). For the therapeutic interventions, the therapeutic modality itself was also specified. For example, within this category there were a number of studies that tested cognitive behavioral therapy (CBT), dialectical behavioral therapy (DBT) and so on.

2.4. Study Quality

An assessment of study quality was conducted. For all RCTs, this was assessed based on the Cochrane Collaboration Risk of Bias Tool [19]. In the majority of trials, as is often the case [24], blinding of participants and therapists was not possible. Each trial was therefore assessed with regard to random sequence generation, allocation concealment, ascertainment of self-harm, outcome assessor blinding, whether analyses were conducted according to the intention-to-treat (ITT) principle, and rates of attrition. For the latter criterion, an attrition rate of 15% or less on the primary outcome at the longest follow-up point indicated low risk of bias.

Non-RCTs were assessed in two ways. For those conducted in clinical, educational, or workplace settings (where a range of study designs were employed) we used a set of criteria based on resources from the Cochrane Effective Practice and Organization of Care (EPOC) group [25]. We assessed whether or not: [1] the study was adequately powered; [2] outcome assessors were blinded to treatment allocation

(for studies where outcomes were measured via interview); [3] the attrition rate was below 15%; and [4] the authors used statistical testing to measure change.

Studies in community settings employed either an ecological or interrupted time series design. Here two criteria were used to assess quality: whether or not data were collected at multiple time points before and after the intervention [26], and whether or not the intervention itself was likely to affect data collection. "Multiple time points" was defined as at least twice before or after implementation of the intervention. The intervention was considered not to affect data collection if sources and methods of data collection were the same before and after the intervention, or if data were collected from official sources (e.g. coronial records).

2.5. Data Synthesis

Meta-analysis was only conducted for RCTs. We analyzed data separately according to study setting. Because self-harm can encompass suicide attempts, is a key predictor of future suicide [27], and is more prevalent and more commonly assessed than suicide, self-harm (measured dichotomously) was our primary outcome and all dichotomous self-harm and suicide attempt data were combined. Additional outcomes were self-harm measured continuously, suicide and suicidal ideation (measured dichotomously and continuously). Where studies had more than one intervention arm, we included those arms that provided relevant data and split the control group to avoid double counting [28].

For dichotomous data, we pooled data between studies using the relative risk with 95% confidence interval. For continuous outcomes, given the range of different tools used, means and standard deviations were pooled using the standardized mean difference (SMD) using the Hedges' adjusted *g* with a 95% confidence interval. SMD effect sizes of 0.2 were considered small, 0.5 were considered medium, and ≥ 0.8 were considered large [29]. Measurement scales were standardized so that higher scores were indicative of greater levels of suicidal ideation. For both continuously- and dichotomously-measured outcomes, pooled effect size estimates were calculated using the DerSimonian-Laird random effects model [30] implemented using Comprehensive Meta-Analysis 2.2.064 software [31].

Between-study heterogeneity was measured using the I^2 statistic. I^2 values of 25%, 50% and 75% or larger are indicative of small, moderate and high heterogeneity, respectively [32].

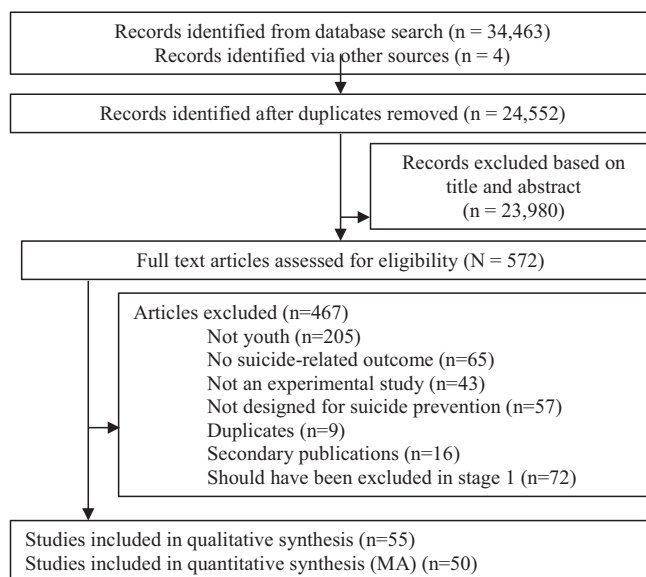


Fig. 1. PRISMA flow diagram.

2.5.1. Subgroup Analysis

For the primary outcome we undertook three subgroup analyses to investigate whether the *intervention approach*, *intervention type* and, for those interventions coded as psychotherapy, the *therapeutic modality* modified the pooled effect sizes.

First, *intervention approach* was coded as universal, selective or indicated. Second, *type* of intervention was categorized as psychotherapy, brief contact, or educational. Psychotherapy interventions were established psychotherapeutic approaches belonging to a particularly theoretical or philosophical school. Brief contact interventions were defined as those interventions that either: [1] focused on maintaining contact or facilitating re-engagement with services via a minimal amount of supportive contact, including provision of an emergency or crisis card as defined by Milner et al. [33]; or [2] interventions delivered within a very brief period, such as screening and referral or provision of one-off assessment and supportive therapy. Educational interventions delivered psycho-education about suicide-related behaviors, mental illness associated with these behaviors, signs and symptoms to look out for and advice on how to respond. Finally, trials coded as psychotherapy were further categorized by *modality* as either: CBT; DBT; mentalisation therapy; problem solving; motivational interviewing; supportive therapy; family therapy; interpersonal psychotherapy; combined (where several modes of psychotherapy were combined); or other (where the intervention did not clearly fit any category of named therapeutic approach).

2.5.2. Sensitivity Analysis

The robustness of results of the meta-analysis was checked for the primary outcome by conducting sensitivity analyses. RCTs judged as high or unclear risk of bias for allocation concealment, and RCTs where more than 15% of participants were lost to follow-up or where no data were reported, were excluded from this analysis.

For studies in which no data amenable to meta-analysis were reported, a narrative synthesis of results was conducted.

3. Results

3.1. Search Results

In total, 34,463 articles were retrieved via database searching and an additional four via the reference lists of included articles. Following initial screening, 572 full-text articles were retrieved, of which 105 met our inclusion criteria. Six were secondary publications that were included as they reported novel data [34–39]. The review therefore includes findings from 105 articles corresponding to 99 unique studies (see Fig. 1).

3.2. Overall Description of Included Studies

Half (52.5%) of included studies were conducted in clinical settings (Tables 1 and 2), 31 (31.3%) in educational or workplace settings (Tables 3 and 4), and 16 (16.2%) in community settings (Tables 5 and 6). Most studies tested indicated interventions ($k = 66$; 66.7%), followed by universal ($k = 17$; 17.2%), multimodal ($k = 11$; 11.1%), and selective ($k = 2$; 2.0%) interventions. Three studies (3.0%) evaluated multiple interventions. Forty-eight studies (48.5%) were RCTs. This included 33 (63.5%) of the studies conducted in clinical settings and 15 (48.4%) of those conducted in educational or workplace settings. None of the community-based studies were RCTs.

The majority of studies were conducted in the United States of America ($k = 49$; 49.5%), followed by the United Kingdom ($k = 12$; 12.1%) and Australia ($k = 11$; 11.1%). Some were conducted across multiple countries and only two (2.0%) were conducted in low-middle income countries. The number of studies more than doubled in the period of 2005–2017 compared to 1990–2004.

3.3. Studies Conducted in Clinical Settings

Fifty-two of the included studies were conducted in clinical settings and all tested indicated interventions delivered to young people with a history of self-harm or attempted suicide resulting in presentation to hospital-based or mental health services. Outcomes therefore refer to repeated self-harm in these studies. Thirty-three were RCTs. Forty (76.9%) had a mean participant age of 18 years or younger, eight studies (15.4%) had a mean age over 18, and in four studies (7.7%) the mean age could not be determined.

3.3.1. Randomized Controlled Trials

3.3.1.1. Study Description. Participants were recruited from emergency departments, inpatient units and community mental health services/outpatient clinics. One study was set in a military hospital [40]. Studies examined the impact of a range of interventions, including individual and group cognitive behavioral therapy (CBT), dialectical behavioral therapy (DBT), family therapy, and brief contact interventions. Control conditions included TAU, e.g. routine care, enhanced TAU, e.g. safety monitoring and facilitated referrals, and active placebo e.g. problem oriented support but without a specific skills-based training component. Twenty-four (72.7%) of the studies in this category included participants with a mean age of 18 or younger. Please see Table 1.

3.3.1.2. Study Efficacy. Thirty-two of the 33 clinical RCTs reported data amenable to meta-analysis. Twenty-five were psychological interventions [40–64] and seven were brief contact interventions [65–71]. The results of the meta-analysis, classified according to outcome assessed, are reported below. The primary outcome (self-harm) is reported first, followed by suicidal ideation; suicide is reported last as it was least frequently assessed.

3.3.1.2.1. Self-harm Measured Dichotomously. Compared to controls, there was no evidence of any intervention effect on self-harm at post-intervention ($k = 12$, $RR = 0.889$, 95% CI 0.71 to 1.11, $I^2 = 37.1\%$) (Fig. 2). At follow-up there was some evidence of a reduction in the proportion of people who had received an intervention who went on to have a repeat self-harm episode ($k = 16$, $RR = 0.83$, 95% CI 0.70 to 0.99, $I^2 = 40.9\%$) (Fig. 3).

3.3.1.2.2. Sensitivity Analysis. There was no material change to the outcome at post-intervention when studies at high risk of bias for allocation concealment were removed. At follow-up, when studies at high risk of bias were removed, the effect was no longer significant.

3.3.1.2.3. Subgroup Analysis. There was no evidence that the *type of intervention* modified the size of the treatment effect post-intervention ($p = 0.67$) or at follow-up ($p = 0.09$); nor was there any evidence that *therapy modality* modified the size of the treatment effect post-intervention ($p = 0.13$), or at follow-up ($p = 0.08$).

3.3.1.2.4. Self-harm Measured Continuously. Compared to controls, there was little evidence, with high heterogeneity ($I^2 = 94.4\%$), that the intervention resulted in a reduction in the mean number of self-harm episodes at post-intervention ($k = 5$, $SMD = -0.66$, 95% CI -1.45 to 0.13), and there was limited evidence of this at follow-up ($k = 4$, $SMD = -0.23$, 95% CI -0.49 to 0.03 , $I^2 = 38.9\%$).

3.3.1.2.5. Suicidal Ideation Measured Dichotomously. Compared to controls, there was no evidence of any effect of intervention on the proportion of people who experienced suicidal ideation post-intervention ($k = 7$, $RR = 0.89$, 95% CI 0.68 to 1.16, $I^2 = 83.0\%$) or at follow-up ($k = 5$; $RR = 0.84$, 95% CI 0.64 to 1.09, $I^2 = 74.8\%$). Heterogeneity was high.

3.3.1.2.6. Suicidal Ideation Measured Continuously. Compared to controls, there was strong evidence of a small effect of the intervention on suicidal ideation post-intervention ($k = 15$, $SMD = -0.28$, 95% CI -0.48 to -0.08 , $I^2 = 76.3\%$). The effect was smaller at follow-up ($k = 11$, $SMD = -0.18$, 95% CI -0.34 to -0.02 , $I^2 = 41.1\%$).

Table 1
Randomized controlled trials conducted in clinical settings (N = 33).

Study; country	Target population	Participants	Intervention description	Comparison condition	Risk of bias	Suicide related outcome(s) assessed; longest follow-up
Alavi et al. (2013) [41] Iran	Inclusion: Young people admitted to hospital for a SA Exclusion: SH w/o intent; no current SI; inability to participate in psychotherapy; diagnosed with bipolar, psychosis, pervasive developmental or substance use disorders Recruited from: Hospital/ED	Whole sample N = 30 Mean age: 16.1 (SD: 1.4; Range: 12–18 Gender: 10% male Treatment group N = 15 Mean age: 16.1 (SD: 1.6) Gender: 6.7% male Control group N = 15 Mean age: 16.0 (SD: 1.2) Gender: 13.3% male	Individual cognitive behavioral therapy plus TAU Length: 12 sessions over 3 months Developed by: Stanley et al. (2009) ^a Delivered by: NR	TAU: routine psychiatric intervention and follow up; pharmacotherapy if needed.	Random sequence generation method: Alternate allocation Allocation concealment method: NR Ascertainment of SH repetition: Self-report Outcome assessor blinding: NA Less than 15% drop-out rate: Yes (0.0%) Was ITT analysis undertaken: NR	SI (continuous): Beck Scale for Suicidal Ideation (BSSI) Longest follow-up: Post-intervention only
Asarnow et al. (2011) [65] USA	Inclusion: Young people who presented to ED with SA or SI Exclusion: Acute psychosis or other symptoms that impede consenting and/or assessment process Recruited from: Hospital/ED	Whole sample N = 181 Mean age: 14.7 (SD: 2.0; Range: 10–18) Gender: 30.9% male Treatment group N = 89 Mean age: 14.8 (SD: 2.1) Gender: 33.7% male Control group N = 92 Mean age: 14.6 (SD: 1.9) Gender: 28.3% male	Brief contact intervention Compliance enhancement measures mixed with family therapy plus TAU Length: 1 month Developed by: Based on Rotheram-Borus et al. (1996) ^b and adapted by authors Delivered by: MH professionals	Enhanced TAU: usual ED care, with staff education on linking to treatment, reducing access to means, risks of substance use.	Random sequence generation method: Computer generated algorithm Allocation concealment method: Independent researcher Ascertainment of SH repetition: Interview Outcome assessor blinding: Yes Less than 15% drop-out rate for SH at post-intervention: Yes (11.6%) Was ITT analysis undertaken: Yes	SI (dichotomous): DISC-IV, an clinician administered diagnostic interview SA (dichotomous): DISC-IV, an clinician administered diagnostic interview and Harkavy Asnis Scale (HASS) Longest follow-up: Post-intervention only
Asarnow et al. (2017) [42] USA	Inclusion: i) Young people who had presented after engaging in SH (SA or NSSI included) within the last three months; ii). history of repetitive SH (≥3 lifetime episodes) Exclusion: symptoms interfering with participation in assessments or intervention (psychosis, substance use) and inability to speak English Recruited from: Hospital/ED and MH outpatient	Whole sample N = 42 Mean age: 14.62 (SD: 1.83) Gender: 11.9% male Treatment group N = 20 Mean age: 14.35 (SD: 1.81) Gender: 10.0% male Control group N = 22 Mean age: 14.86 (SD: 1.86) Gender: 13.6% male	SAFETY program Combined intervention consisting of CBT and DBT informed family intervention that included formulation driven CBT, DBT and family centered interventions. Each family had two therapists: one for the young person and one for the parents and there were joint family sessions as well as separated sessions. Length: 12 sessions over 3 months Developed by: study authors Delivered by: MH professionals	Enhanced TAU: in-clinic parent education on risk of repetition, accessing treatment; 3 + phone-calls monitoring safety, encouraging treatment attendance.	Random sequence generation method: Computerized randomization program Allocation concealment method: Enrolment and assessment staff masked to randomization status Ascertainment of DSH repetition: Self-report Outcome assessor blinding: Yes Less than 15% drop-out rate: No (23.8% for self-reported outcomes) Was ITT analysis undertaken: Yes	SA (dichotomous): used a slight modification of the clinician administered Columbia Suicide Severity Rating Scale (C-SSRS) Longest follow-up: 12 months post-baseline
Bertolote et al. (2010) [66]; Fleischmann et al. (2008) [34] Multi-national	Inclusion: Young people who presented to ED following SH/self-poisoning Exclusion: 'any clinical condition (s) that would disallow interview' Recruited from: Hospital/ED	Whole sample N = 1867 Mean age: NR (Median = 23.0) Gender: 41.8% male Treatment group N = 922 Mean age: NR Gender: 40% male Control group N = 945 Mean age: NR Gender: 43.3% male	Brief contact intervention 1 1-hour information session plus 9 phone calls or visits. Length: Up to 10 contacts over 18 months Developed by: study authors (based on existing BIC methods) Delivered by: doctor, nurse or psychologist	TAU: varied between sites, primarily acute injury management with or without mental health referral.	Random sequence generation method: Random numbers table Allocation concealment method: Offsite researcher Ascertainment of SH repetition: Interview Outcome assessor blinding: NR Less than 15% drop-out rate: Yes (11.0%) Was ITT analysis undertaken: NR	SA (dichotomous): European Parasuicide Study Interview Schedule (EPSIS) of the WHO/EURO Multicenter Study on Suicidal Behavior Longest follow-up: Post-intervention only

Byford et al. (1999) [43] UK	Inclusion: Diagnosis of SH (self-poisoning) Exclusion: Overdose was accidental; psychiatric condition which would preclude engagement with therapy; social situation precluded engagement with family therapy Recruited from: MH outpatient	Whole sample N = 162 Age/gender: NR Treatment group N = 85 Age/gender: NR Control group N = 77 Age/gender: NR	Individual family therapy plus TAU Length: 1 ½ hour assessment plus 1 h of therapy Developed by: study authors Delivered by: MH professionals	TAU: routine assessment and psychiatric care in outpatient clinic.	Random sequence generation method: Shuffled cards Allocation concealment method: Sealed envelope Ascertainment of SH repetition: interview Outcome assessor blinding: Yes Less than 15% drop-out rate: Yes (8.0%) Was ITT analysis undertaken: NR	SI (continuous): Suicidal Ideation Questionnaire (SIQ) Longest follow-up: Post-intervention only
Carter et al. (2010) [44] Australia	Inclusion: Females referred for treatment following self-poisoning, meeting criteria for borderline personality disorder, with at least three self-reported episodes of self-harm over the preceding year. Exclusion: Males, those engaging in self-injury without self-poisoning Recruited from: MH outpatient	Whole sample N = 70 Mean age: 24.5 (SD: 6.1; Range: 18–65) Gender: 0% male Treatment group N = 37 Mean age: 24.5 (SD: 6.1) Control group N = 33 Mean age: 24.7 (SD: 6.2)	Dialectical behavior therapy Individual and group therapy, with telephone coaching. Length: number of sessions not specified, delivered over six months Developed by: based on Linehan et al. (1991) ^c Delivered by: MH professionals	TAU + Waitlist: 6 month period of unspecified TAU while waitlisted.	Random sequence generation method: Shuffled envelopes Allocation concealment method: Sealed, opaque envelopes Ascertainment of SH repetition: Interview Outcome assessor blinding: Yes Less than 15% drop-out rate: Yes (0.0%) Was ITT analysis undertaken: Mixed methods	SH (continuous): Linehan's Lifetime Parasuicide Count–2; Parasuicide History Interview SH (dichotomous): Linehan's Lifetime Parasuicide Count–2; Parasuicide History Interview Longest follow-up: Post-intervention only
Cooney et al. (2010) [45] New Zealand	Inclusion: History of at least one SA or one episode of SH in past three months Exclusion: i) Intellectual disability; ii) Psychosis Recruited from: MH outpatient	Whole sample N = 29 Mean age: 15.9 (SD: 1.0; Range: 14–18) Gender: 24.1% male Treatment group N = 14 Mean age: 16.2 (SD: 0.98) Gender: 28.6% male Control group N = 15 Mean age: 15.7 (SD: 1.1) Gender: 20% male	Individual plus group dialectical behavioral therapy Length: weekly sessions for approximately 26 weeks Developed by: based on Linehan (1993) ^d & Miller et al. (2007) ^e Delivered by: MH professional	TAU: type and duration varied: CBT, motivational interviewing, supportive counseling, family therapy; medication and case management as needed.	Random sequence generation method: Computer generated algorithm Allocation concealment method: Sealed, opaque envelopes Ascertainment of SH repetition: Interview Outcome assessor blinding: Yes Less than 15% drop-out rate: Yes (0.0%) Was ITT analysis undertaken: Mixed methods	SI (continuous): BSSI SA (dichotomous): Linehan's Suicide Attempt-Self-Injury Interview (SASII) Longest follow-up: Post-intervention only
Cotgrove et al. (1995) [67] UK	Inclusion: Admitted to hospital following SA/SH Exclusion: Records of the original SA were missing, or were there insufficient follow-up data (p. 572) Recruited from: Hospital/ED	Whole sample N = 105 Mean age: 14.9 (SD: NR; Range: 12.2–16.7) Gender: 15.2% male Treatment group N = 4 Age/gender: NR Control group N = 58 Age/gender: NR	Brief contact intervention Emergency card allowing readmission to hospital on request. Length: NA Developed by: based on Morgan et al. (1993) ^f Delivered by: NA	TAU: standard follow-up care per ED site.	Random sequence generation method: Open random numbers table Allocation concealment method: Use of an open random numbers table suggests allocation could not have been concealed Ascertainment of SH repetition: Hospital and clinical notes Outcome assessor blinding: NA Less than 15% drop-out rate: Yes (0.0%) Was ITT analysis undertaken: NR	SA (dichotomous): information collected from clinic and hospital records, and contacting general practitioners and other health professionals involved with young person Longest follow-up: 12 months post-baseline
Diamond et al. (2010) [46] ^g	Inclusion: i) Scored >31 on the SIQ-JR (Reynolds, 1987) ^h ; ii) score remained elevated 2 days later	Whole sample N = 66 Mean age: 15.2 (SD: 1.62; Range 12–17)	Individual family therapy plus TAU Length: Up to 15 sessions	Enhanced TAU: safety monitoring and facilitated referrals for treatment (incl.	Random sequence generation method: Adaptive randomization	SI (continuous): SIQ-Junior (SIQ-JR) SI (dichotomous): SIQ-JR

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Table 1 (continued)

Study; country	Target population	Participants	Intervention description	Comparison condition	Risk of bias	Suicide related outcome(s) assessed; longest follow-up
USA	following a second screen. Exclusion: i) Current psychosis; ii) mental retardation/history of borderline intellectual functioning Recruited from: Hospital/ED and primary care practices (75.0% were recruited from primary care and 25.0% from hospitals/EDs)	Gender: 16.7% male Treatment group N = 35 Mean age: 15.1 (SD: 1.41) Gender: 8.6% male Control group N = 31 Mean age: 15.3 (SD: 1.83) Gender: 25.8% male	delivered over a 3-month period Developed by: study authors Delivered by: trained PhD or Masters level therapists	Individual, group, or family therapy, or case management).	Allocation concealment method: Independent researcher Ascertainment of SH repetition: Interview Outcome assessor blinding: No Less than 15% drop-out rate: Yes (0.0%) Was ITT analysis undertaken: Yes	Longest follow-up: 3 months post-intervention
Donaldson et al. (2005) [47] USA	Inclusion: Presented to general pediatric child psychiatric hospital after SA. Exclusion: Current psychosis Recruited from: Hospital/ED	Whole sample N = 39 Mean age: 15.0 (SD: 1.7; Range: 12–17) Gender: 18% male Treatment group: N = NR Age/gender: NR Control group: N = NR Age/gender: NR	Individual skills-based therapy ⁱ by trained therapists Length: 12 sessions delivered over 6 months Developed by: Study authors Delivered by: Trained therapists	Enhanced TAU: Supportive Relationship Treatment (SRT).	Random sequence generation method: Random numbers table Allocation concealment method: NR Ascertainment of SH repetition: Interview Outcome assessor blinding: NR Less than 15% drop-out rate: No (20.5%) Was ITT analysis undertaken: No	SI (continuous): SIQ SA (dichotomous): Structured adolescent follow-up interviews Longest follow-up: 6 months post-baseline
Esposito- Smythers et al. (2011) [48] USA	Inclusion: SA in past 3 months or scored >41 on the SIQ (Reynolds, 1987) Exclusion: Verbal IQ score < 70; ii) Psychosis; iv) Bipolar disorder; iv) Dependent on substances other than alcohol or cannabis Recruited from: MH outpatient	Whole sample N = 40 Mean age: 15.7 (SD: 1.19; Range: 13–17) Gender: 33.3% male Treatment group N = 20 Mean age: 15.8 (SD: 0.98) Gender: 31.6% male Control group N = 20 Mean age: 15.7 (SD: 1.41) Gender: 35.3% male	Individual cognitive behavioral therapy Length: 24 sessions delivered over 12 months Developed by: based on Donaldson et al. (2005) and Esposito Smythers et al. (2006) and adapted by study authors Delivered by: Trained therapists	Enhanced TAU: treatment schedule and approach determined by community providers. Diagnostic evaluation report provided. Study psychiatrist assisted with medication management. Access to information and resources.	Random sequence generation method: Computer generated adaptive randomization Allocation concealment method: Unclear Ascertainment of SH repetition: Interview Outcome assessor blinding: Assessors could guess allocation due to offhand comments made by participants during interviews Less than 15% drop-out rate: No (25.0%) Was ITT analysis undertaken: Yes	SI (continuous): SIQ SA (dichotomous): Kiddie Schedule for Affective Disorders and Schizophrenia (K-SADS) – clinician administered diagnostic interview. Longest follow-up: 6 months post-intervention
Green et al. (2011) [49] UK	Inclusion: Presented to child and adolescent services with at least two episodes of SH in the past 12 months Exclusion: i) Severe low weight anorexia nervosa; ii) psychosis; iii) learning disability Recruited from: MH outpatient	Whole sample N = 366 Mean age: NR (Range: 12–16) Gender: 11.5% male Treatment group N = 183 Mean age: NR Gender: 11.5% male Control group N = 183 Mean age: NR Gender: 11.5% male	Group cognitive behavioral therapy Length: 6 sessions during the acute phase & as many sessions needed during the maintenance phase Developed by: based on Wood et al. (2001) Delivered by: Trained therapists	TAU: routine care provided by local child & adolescent mental health services according to clinical judgment, excluding group interventions.	Random sequence generation method: Computer generated minimization algorithm Allocation concealment method: Independent, off-site researcher Ascertainment of SH repetition: Interview Outcome assessor blinding: Yes Less than 15% drop-out rate: Yes (4.0%) Was ITT analysis undertaken: No	SI (continuous): SIQ-JR SI (dichotomous): SIQ-JR SH (dichotomous): SIQ-JR Longest follow-up: 12 months post-baseline
Harrington et al. (1998) [50] ^j UK	Inclusion: Presented to hospital with self-poisoning Exclusion: i) Other SH (e.g.	Whole sample N = 162 Mean age: 14.5 (SD: 1.15; Range: 10–16)	Five sessions of family therapy plus TAU Length: NR	TAU: routine psychiatric aftercare including diverse range of interventions, but no	Random sequence generation method: Shuffled envelopes Allocation concealment	SI (continuous): SIQ SI (dichotomous): SIQ Suicide: NR

	cutting); ii) Severe suicidality; iii) clinician determined risk of contraindication for family treatment, e.g. psychosis, currently receiving psychiatric treatment, parent/child had a learning difficulty	Gender: 10.5% male Treatment group N = 85 Mean age: 14.4 (SD: 1.2) Gender: 10.6% male Control group N = 77 Mean age: 14.6 (SD: 1.1) Gender: 10.6% male	Developed by: study authors Delivered by: 2 experienced masters'-level child psychiatric social workers	home-based family interventions.	method: Sealed, opaque envelopes Ascertainment of SH repetition: Interview Outcome assessor blinding: Attempted but not always possible Less than 15% drop-out rate: Yes (8.0%) Was ITT analysis undertaken: No	Longest follow-up: 12 months post-baseline
Hassanian-- Moghaddam et al. (2011) [68] Iran	Inclusion: Presented to hospital with self-poisoning Exclusion: Psychosis Recruited from: Hospital/ED	Whole sample N = 2133 Mean age: 24.1 (SD: 8.11; Range: NR) Gender: 33.7% male Treatment group N = 1043 Mean age: 24.7 (SD: 7.97) Gender: 33.3% male Control group N = 1070 Mean age: 24.1 (SD: 8.25) Gender: 34% male	Brief contact intervention (<i>Postcards from Persia</i>) plus TAU. Length: 8 postcards mailed over 12 months Developed by: based on Carter et al. (2005) ^k Delivered by: NA	TAU: follow-up care for self-poisoning in Tehran is "poor", contact is mainly hospital- or office-based.	Random sequence generation method: Block randomization using a random numbers table Allocation concealment method: Allocation was concealed, but information on the method used was not provided Ascertainment of SH repetition: Interview Outcome assessor blinding: No Less than 15% drop-out rate: Yes (8.1%) Was ITT analysis undertaken: No	SI (continuous): follow-up interview SI (dichotomous): follow-up interview SH (continuous): follow-up interview SH (dichotomous): follow-up interview SA (continuous): follow-up interview SA (dichotomous): follow-up interview Suicide: mortality records Longest follow-up: 12 months post-baseline
Hazell et al. (2009) [51] Australia	Inclusion: Presented to hospital with >2 episodes of SH Exclusion: i) Acute psychosis; ii) intellectual disability Recruited from: MH outpatient	Whole sample N = 72 Mean age: 14.5 (SD: 1.1; Range 12–16) Gender: 9.7% male Treatment group N = 35 Mean age: 14.6 (SD: 1.1) Gender: 8.6% male Control group N = 37 Mean age: 14.4 (SD: 1.2) Gender: 10.8% male	Group based cognitive behavioral therapy (<i>Moving on from self-harm</i>) plus TAU Length: 6 sessions over 12 months Developed by: study authors Delivered by: MH professionals	TAU: routine care varied but generally included individual/family counseling, medication assessment, and care-coordination.	Random sequence generation method: Block randomization using a computer generated random numbers table Allocation concealment method: Independent, offsite researcher Ascertainment of SH repetition: Interview Outcome assessor blinding: Yes Less than 15% drop-out rate: Yes (0.0%) Was ITT analysis undertaken: Mixed methods	SI (continuous): SIQ SH (dichotomous): Linehan's Parasuicide History Interview Longest follow-up: 12 months post-baseline
Huey et al. (2004) [52] USA	Inclusion: Presented to hospital with SA/SI Exclusion: Autism spectrum disorder Recruited from: Hospital/ED	Whole sample N = 156 Mean age: 12.9 (SD: 2.1; Range 10–17) Gender: 35% male Treatment group N = Unclear Age/gender: NR Control group N = Unclear Age/gender: NR	Multi-systematic family therapy Length: Unclear Developed by: Henggeler et al. (2002) ^l Delivered by: MH professionals	Active placebo: hospitalization at youth inpatient psychiatric unit.	Random sequence generation method: NR Allocation concealment method: NR Ascertainment of SH repetition: Interview Outcome assessor blinding: NA Less than 15% drop-out rate: Unclear Was ITT analysis undertaken: NR	SI (dichotomous): Youth Risk Behavior Survey (YRBS) SA (dichotomous): Child Behavior Checklist (CBCL) Longest follow-up: 12 months post-intervention
Husain et al. (2014) [53] Pakistan	Inclusion: Admitted to hospital following SH Exclusion: i) dementia; ii) substance misuse; iii) organic mental disorder; iv) delirium; v)	Whole sample N = 221 Mean age: 23.1 (SD: 5.5; Range: 16–64) Gender: 31.2% male Treatment group	Individual cognitive behavioral therapy (<i>Life After Self-harm</i>) plus TAU Length: 6 sessions over 3 months Developed by: based on Schmidt	TAU: standard routine care provided by local services.	Random sequence generation method: Computer generated random numbers table Allocation concealment method: Independent, offsite	SI (continuous): BSSI Suicide: Not stated Longest follow-up: 6 months post-baseline

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Table 1 (continued)

Study; country	Target population	Participants	Intervention description	Comparison condition	Risk of bias	Suicide related outcome(s) assessed; longest follow-up
	alcohol and/or drug dependence; vi) schizophrenia; vii) bipolar disorder; viii) intellectual disability Recruited from: Hospital/ED	N = 108 Mean age: 23.2 (SD: 5.8) Gender: 29.6% male Control group N = 113 Mean age: 23.1 (SD: 5.3) Gender: 32.7%	& Davidson (2004) ^m and adapted by study authors Delivered by: masters-level psychologists		researcher Ascertainment of SH repetition: Interview Outcome assessor blinding: Yes Less than 15% drop-out rate: Yes (3.6% by the 6-month follow-up period; could not calculate for final follow-up) Was ITT analysis undertaken: Yes	
King et al. (2006) [54] USA	Inclusion: i) SA or severe SI in past 3 months ii) Score of 20 or 30 on the Self-Harm subscale of the Child and Adolescent Functional Assessment Scale (Hodges, 1989) ⁿ Exclusion: i) Severe intellectual disability; ii) Psychosis Recruited from: Hospital/ED	Whole sample N = 289 Mean age: 15.3 (SD: 1.5; Range: 12–17) Gender: 31.8% male Treatment group N = 151 Mean age: 15.4 (SD: 1.5) Gender: 31.1% male Control group N = 138 Mean age: 15.2 (SD: 1.4) Gender: 32.6% male	Supportive intervention Youth nominated support team Version 1 plus TAU One-off brief psycho-education intervention for support team plus up to 9 contacts per week between adolescent and support team Length: 1.5 to 2 h Developed by: study authors Delivered by: MH professional	TAU: varied, included psychotherapy, medication, alcohol/drug treatment, partial hospitalization, and community services.	Random sequence generation method: Random numbers table Allocation concealment method: Independent researcher Ascertainment of SH repetition: Interview Outcome assessor blinding: No Less than 15% drop-out rate: Yes (18.3%) Was ITT analysis undertaken: Yes	SI (continuous); SIQ-JR SI (dichotomous): SIQ-JR SA (dichotomous): Not stated Longest follow-up: 6 months post-baseline
King et al. (2009) [55] USA	Inclusion: SA or severe SI in past 4 weeks Exclusion: NR Recruited from: Hospital/ED	Whole sample N = 448 Mean age: 15.6 (SD: 1.31; Range: 13–17) Gender: 28.8% male Treatment group N = 223 Mean age: 15.6 (SD: 1.24) Gender: NR Control group N = 225 Mean age: 15.6 (SD: 1.37) Gender: NR	Supportive intervention Youth nominated support team Version 2 plus TAU One-off, individual or group-based (as preferred) psycho-education session plus weekly telephone contacts. For adolescents: weekly sessions by telephone or face-to-face as preferred with support team. Length: 1 h Developed by: study authors Delivered by: MH professional	TAU: as above.	Random sequence generation method: Block randomization using a computer generated sequence Allocation concealment method: Independent researcher Ascertainment of SH repetition: Interview Outcome assessor blinding: Yes Less than 15% drop-out rate for: No (23.0%) Was ITT analysis undertaken: Mixed methods	Suicide: Not stated SA (dichotomous): Clinician administered diagnostic interview DISC-IV Mood Disorders module Longest follow-up: 12 months post-baseline
King et al. (2015) [56] USA	Inclusion: Presented to ED with SI, a recent SA or positive screens for both depression plus alcohol/drug abuse Exclusion: Required referral for inpatient psychiatric hospitalization Recruited from: Hospital/ED	Whole sample N = 49 Mean age: 17.7 (SD: 1.7; Range: 14–19) Gender: 40% male Treatment group N = 27 Age/gender: NR Control group N = 22 Age/gender: NR	Individual motivational interview plus TAU Length: 35–45 min Developed by: study authors (based on standard motivational interviewing protocols) Delivered by: trained therapists	Enhanced TAU: adolescents given a crisis card and written information about depression, suicide, firearm safety, and services.	Random sequence generation method: Shuffled envelopes Allocation concealment method: NR Ascertainment of SH repetition: Interview Outcome assessor blinding: Yes Less than 15% drop-out rate: Yes (6.1%) Was ITT analysis undertaken: Yes	SI (continuous): SIQ-JR Longest follow-up: 2 months post-baseline
McLeavey et al. (1994) [57] Republic of Ireland	Inclusion: Presented to ED with self-poisoning Exclusion: i) Required psychiatric inpatient/day-hospital admission;	Whole sample N = 39 Mean age: 24.4 (SD: 7.0; Range 15–45) Gender: 25.6% male	Individual Interpersonal Problem-Solving Skills Training Length: Five weekly one-hour sessions for 5 weeks (with 1	Active placebo: brief problem-oriented approach, did not involve skills training.	Random sequence generation method: NR Allocation concealment method: NR	SH (dichotomous): ED readmission Suicide: Hospital records

	ii) Psychosis; iii) Intellectual disability; iv) organic cognitive impairment Recruited from: Hospital/ED	Treatment group N = 19 Mean age: 23.6 (SD: 5.9) Gender: 21% male Control group N = 20 Mean age: 25.3 (SD: 8.1) Gender: 30% male	additional session if necessary) Developed by: study authors Delivered by: MH professionals		Ascertainment of SH repetition: GP questionnaire and hospital records Outcome assessor blinding: NA Less than 15% drop-out rate: No (15.4%) Was ITT analysis undertaken: Not described	Longest follow-up: 12 months post-intervention
Mehlum et al. (2016) [58] ^o Norway	Inclusion: Referred to child & adolescent psychiatric outpatient clinic with a history of >2 episodes of self-harm; 1 within the past 16 weeks Exclusion: i) Bipolar disorder (except bipolar II); ii) Schizophrenia; iii) Affective disorder; iv) Psychosis NOS; v) Intellectual disability; vi) Asperger's syndrome Recruited from: MH outpatient & Hospital/ED	Whole sample N = 77 Mean age: 15.6 (SD: 1.6; Range: 12-18) Gender: 11.7% male Treatment group N = 39 Mean age: 15.9 (SD: 1.4) Gender: 12.8% male Control group N = 38 Mean age: 15.3 (SD: 1.6) Gender: 20.5% male	Individual and group Dialectical Behavior Therapy Length: 19 weeks – One 1-hour weekly session of individual therapy; one 2-hour weekly session of multifamily skills training; plus family therapy & telephone coaching as needed. Developed by: Miller et al. (2007) Delivered by: MH professionals	Enhanced TAU: standard care enhanced for the purpose of the trial by requiring that therapists agree to provide at least 1 weekly treatment session per patient.	Random sequence generation method: Block randomization using a computer generated sequence Allocation concealment method: Independent researcher Ascertainment of SH repetition: Interview supplemented with hospital records Outcome assessor blinding: NA Less than 15% drop-out rate: Yes (2.6% by the one-year follow-up period) Was ITT analysis undertaken: Yes	SI (continuous): SIQ-JR SH (continuous): ED readmission and self-report SH (dichotomous): ED readmission and self-report Suicide: Mortality records Longest follow-up: 12 months post-intervention
Ougrin et al. (2011) [69]; (2013) [39] UK	Inclusion: Referred to ED following SH Exclusion: i) Psychosis; ii) Intoxication; iii) Learning disability; iv) Required inpatient admission Recruited from: Hospital/ED	Whole sample N = 70 Mean age: 15.5 (SD: 1.3; Range: 12-18) Gender: 20% male Treatment group N = 35 Mean age: 15.6 (SD: 1.5) Gender: 20% male Control group N = 35 Mean age: 15.5 (SD: 1.2) Gender: 20% male	Brief contact intervention Comprised psychosocial history & risk assessment plus brief intervention Length: 1 h plus 30 min Developed by: study authors Delivered by: MH professionals	TAU: standard psychosocial history and risk assessment, report sent to relevant community team	Random sequence generation method: Block randomization using a computer generated sequence Allocation concealment method: Independent researcher Ascertainment of SH repetition: Interview supplemented with clinical records Outcome assessor blinding: NA Less than 15% drop-out rate: Yes (1.4% by the two-year follow-up period) Was ITT analysis undertaken: Yes	SH (dichotomous): ED readmission Longest follow-up: 24 months post-baseline
Pineda & Dadds, (2013) [59] Australia	Inclusion: Presented to ED with either SI, SA or SH within the 2 months prior to presentation Exclusion: 1) Overdose of recreational drugs; ii) Intellectual disability Recruited from: ED	Whole sample N = 48 Mean age: 15.1 (SD: 1.2; Range: 12-17) Gender: 25% male Treatment group N = 24 Mean age: 15.0 (SD: 1.31) Gender: 27.3% male Control group N = 24 Mean age: 15.28 (SD: 1.18) Gender: 22.2% male	Strengths-based family education program plus TAU: <i>Resourceful Adolescent Parent Program (RAP-P)</i> Length: Four 2-hour sessions delivered in a single family format either once a week or once every two weeks. A total of five, 2-hour sessions were provided over up to 2.5 months. Developed by: based on Shochet et al. (1997) ^p and adapted by study authors Delivered by: primary author	TAU: routine care (included any intervention deemed necessary by the treating team other than RAP-P).	Random sequence generation method: Random numbers table Allocation concealment method: Independent researcher Ascertainment of SH repetition: Interview Outcome assessor blinding: Yes Less than 15% drop-out rate: No (16.7%) Was ITT analysis undertaken: Yes	SI (continuous): ASQ-R Longest follow-up: 6 months post-baseline

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Table 1 (continued)

Study; country	Target population	Participants	Intervention description	Comparison condition	Risk of bias	Suicide related outcome(s) assessed; longest follow-up
Power et al. (2003) [60] Australia	Inclusion: Referred to a specialist first episode psychosis clinic with SI or SA Exclusion: NR Recruited from: MH outpatient	Whole sample N = 56 Age/gender: NR Treatment group N = 31 Age/gender: NR Control group N = 25 Age/gender: NR	Individual cognitive oriented therapy (<i>Lifespan</i>) plus TAU Length: Eight to ten sessions over 10 weeks. Developed by: study authors Delivered by: MH professionals	TAU: standard clinical care.	Random sequence generation method: NR Allocation concealment method: NR Ascertainment of SH repetition: Clinical records Outcome assessor blinding: NA Less than 15% drop-out rate: No (37.5%) Was ITT analysis undertaken: NR	Suicide: Not stated Longest follow-up: Post-intervention only
Robinson et al. (2012) [70] Australia	Inclusion: Referred but not accepted to a specialist outpatient adolescent MH service with a history of SI, SA or SH Exclusion: i) Intellectual disability; ii) Known organic cause for presentation Recruited from: MH outpatient	Whole sample N = 164 Mean age: 18.6 (SD: NR; Range: 15–24) Gender: 35.4% male Treatment group N = 81 Mean age: NR Gender: 39.5% male Control group N = 83 Mean age: NR Gender: 31.3% male	Brief contact intervention plus TAU – monthly postcards Length: Twelve postcards over 12 months Developed by: study authors (based on existing BIC methods) Delivered by: NA	TAU: treatment or support already being received; e.g., from school counselor, GP, psychologist.	Random sequence generation method: Block randomization using a computer generated sequence Allocation concealment method: Independent researcher Ascertainment of SH repetition: Interview Outcome assessor blinding: Yes Less than 15% drop-out rate: No (52.7%) Was ITT analysis undertaken: No. However, sensitivity analyses were undertaken which suggested that ITT results with data imputed for all missing observations not materially different to per protocol analysis	SI (continuous): BSSI SI (dichotomous): BSSI SH (dichotomous): Suicide Behavior Questionnaire-14 item version (SBQ-14) SA (dichotomous): SBQ-14 Longest follow-up: 6 months post-intervention
Rossouw & Fonagy, (2012) [61] UK	Inclusion: Presented to ED or referred to community MH services with SH Exclusion: i) Presentation the result of excessive use of recreational drugs; ii) Psychosis; iii) Severe learning disability; iv) Developmental disorder; v) Eating disorder; vi) Dependence on alcohol/drugs Recruited from: Hospital/ED and MH outpatient	Whole sample N = 80 Mean age: 14.7 (SD: 1.25; Range: 12–17) Gender: 15% male Treatment group N = 40 Mean age: 15.4 (SD: 1.3) Gender: 17.5% male Control group N = 40 Mean age: 14.8 (SD: 1.2) Gender: 12.5% male	Mentalization therapy: comprised weekly individual sessions plus monthly family therapy. Length: 1 year Developed by: study authors Delivered by: MH professionals.	TAU: routine care provided by community-based adolescent mental health services. Mainly individual therapeutic intervention, combined individual and family therapy, or psychiatric review.	Random sequence generation method: Minimization algorithm Allocation concealment method: Independent, offsite researcher Ascertainment of SH repetition: Interview Outcome assessor blinding: Yes Less than 15% drop-out rate: Yes (11.2%) Was ITT analysis undertaken: Yes	Suicide: Not stated SH (continuous): Risk-Taking and Self-Harm Inventory (RTSHI) SH (dichotomous): RTSHI Longest follow-up: Post-intervention only
Rudd et al. (1996) [40] ^a USA	Inclusion: Referred to outpatient mental health clinics, an inpatient service or an ED with SA, SI Exclusion: i) Substance abuse/dependence ii) Psychosis/thought disorder; iii) Personality disorder Recruited from: Hospital/ED and	Whole sample N = 264 Mean age: 22.2 (SD: 2.3; Range: NR) Gender: 82.2% male Treatment group N = 143 Mean age: NR Gender: 77.6% male	Group-based problem-solving and social competence training Length: 9 h a day for two weeks Developed by: study authors Delivered by: MH professionals	TAU: combination of inpatient and outpatient care.	Random sequence generation method: Sequential randomization Allocation concealment method: NR Ascertainment of SH repetition: Interview Outcome assessor blinding: NR	SI (continuous): Modified Scale for Suicidal Ideation (MSSI) ^f Longest follow-up: Post-intervention only

	MH outpatient NB: Setting comprised ¹ y medical center	Control group N = 121 Mean age: NR Gender: 87.6% male			Less than 15% drop-out rate: No (73.1% by the 12 month follow-up period) Was ITT analysis undertaken: Unclear	
Slee et al. (2008) [62] ^s The Netherlands	Inclusion: Presented to an outpatient MH service with recent SH Exclusion: Psychiatric disorder requiring inpatient treatment Recruited from: MH outpatient	Whole sample N = 82 Mean age: 24.6 (SD: 5.4; Range: 15-35) Gender: 6.1% male Treatment group N = 40 Mean age: 23.9 (SD: 6.4) Gender: 2.5% male Control group N = 42 Mean age: 25.4 (SD: 4.5) Gender: 9.5% male	12 out-patient, individual cognitive behavioral therapy sessions plus TAU Length: weekly sessions for up to 5.5 months Developed by: NR (but based on standard CBT protocol) Delivered by: MH professionals	TAU: participants' choice, three forms: psychotropic medication, psychotherapy and psychiatric hospitalizations	Random sequence generation method: Computer generated random numbers table Allocation concealment method: Independent, offsite researcher Ascertainment of SH repetition: Interview Outcome assessor blinding: NR. Less than 15% drop-out rate: No (21.0%) Was ITT analysis undertaken: Mixed methods	SH (continuous): Structured clinical interview SH (dichotomous): Structured clinical interview Longest follow-up: 9 months post-baseline
Spirito et al. (2002) [71] USA	Inclusion: Presented to an ED/pediatric hospital with SA Exclusion: NR Recruited from: Hospital/ED	Whole sample N = 63 Mean age: 15.0 (1.4; Range: 12-18) Gender: 9.5% male Treatment group N = 29 Mean age: NR Gender: 13.8% male Control group N = 34 Mean age: NR Gender: 5.9% male	Brief contact intervention Individual compliance enhancement intervention plus TAU Length: one hour Developed by: study authors Delivered by: post-doctoral psychology fellows	TAU: standard disposition planning.	Random sequence generation method: Random numbers table Allocation concealment method: NR Ascertainment of SH repetition: Interview Outcome assessor blinding: NR Less than 15% drop-out rate: No (17.1%) Was ITT analysis undertaken: No	SH (dichotomous): Structured interview Suicide: Not stated Longest follow-up: 3 months post-baseline
Spirito et al. (2015) [72] USA	Inclusion: Resided in a specific catchment plus current or past 'suicidality' Exclusion: NR Recruited from: MH outpatient and Hospital/ED	Whole sample N = 24 Mean age: 14.3 (SD: 1.7; Range: 11-17) Gender: 16.7% male Treatment group N = 16 Mean age: 14.7 (SD: 1.8) Gender: 12.5% male Control group N = 8 Mean age: 14.0 (SD: 1.7) Gender: 25% male	Parent-Adolescent-cognitive behavioral therapy Individual CBT (for the parents plus adolescents) and family sessions Length: 12 sessions over 12 weeks Developed by: based on protocols used in prior clinical trials with depressed Adolescents. Delivered by: masters and PhD level clinicians	Active placebo: adolescent-only CBT.	Random sequence generation method: NR Allocation concealment method: NR Ascertainment of DSH repetition: Interview Outcome assessor blinding: NR Less than 15% drop-out rate: Unclear Was ITT analysis undertaken: Yes	SI (continuous): BSSI-A Longest follow-up: 12 months post-baseline Not included in MA ²
Wharff et al. (2017) [63] USA	Inclusion: i) presentation to ED with "suicidality" or suicide attempt; ii) presence of consenting parent or legal guardian Exclusion: i) not fluent in English; ii) Not medically stable, including intoxication; iii) cognitive 'limitations' preventing completion of research instruments; iv) active psychosis; v) required physical or medical restraint in ED Recruited from: Hospital/ED	Whole sample N = 142 Mean age: 15.5 (SD: 1.4) Gender: 28% male Treatment group N = 68 Mean age: 15.4 (SD: 1.3) Gender: 26% male Control group N = 71 Mean age: 15.6 (SD: 1.5) Gender: 30% male	Family Based Crisis Intervention (based on cognitive behavioral therapy) plus TAU; an emergency crisis intervention Length: 60 to 90-min Developed by: study authors Delivered by: master's level psychiatric social workers	TAU: standard psychiatric Evaluation and clinical/discharge recommendations.	Random sequence generation method: NR Allocation concealment method: NR Ascertainment of DSH repetition: NA Outcome assessor blinding: NA Less than 15% drop-out rate: Yes (19.0%) Was ITT analysis undertaken: No	SI (continuous): Reasons for Living Inventory for Adolescents (RFL-A) ^t Longest follow-up: 1 month post-intervention
Wood et al. (2001) [64]	Inclusion: i) Referred to child & adolescent MH service following	Whole sample N = 63	Combined group-based psychotherapy plus TAU.	TAU: variety of interventions delivered by community	Random sequence generation method: Random numbers	SH (continuous): ED readmission (assessed via

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Table 1 (continued)

Study; country	Target population	Participants	Intervention description	Comparison condition	Risk of bias	Suicide related outcome(s) assessed; longest follow-up
UK	SH; ii) Engaged in SH on at least one other occasion during the past year Exclusion: i) 'Too suicidal' for ambulatory care; ii) psychosis; iii) learning 'problems' Recruited from: MH outpatient	Mean age: 14.3 (SD: 1.6; Range: 12–16) Gender: 22.2% male Treatment group N = 32 Mean age: 14.2 (SD: 1.1) Gender: 21.9% male Control group N = 31 Mean age: 14.3 (SD: 2.1) Gender: 25.8% male	Comprised aspects of cognitive behavioral therapy, dialectical behavioral therapy and psychodynamic psychotherapy. Length: "until the young person feels ready to leave" (p. 1247). Developed by: study authors Delivered by: MH professionals	psychiatric nurses & psychologists. Included family sessions, nonspecific counseling. Psychotropic medication (where indicated).	table Allocation concealment method: Independent, offsite researcher Ascertainment of SH repetition: Interview Outcome assessor blinding: Yes Less than 15% drop-out rate: Yes (3.1%) Was ITT analysis undertaken: Yes	interview) SH (dichotomous): ED readmission Suicide: Not stated Longest follow-up: 7 months post-randomization

Notes: ED = Emergency Department; ITT = intention-to-treat; IQR = Interquartile Range; MA = meta-analysis; MH = mental health; NR = not reported; TAU = treatment as usual; SA = suicide attempt; SD = standard deviation; SH = self-harm; SI = suicidal ideation; SRB = suicide-related behavior.

- ^a Stanley B, et al. Cognitive behavioral therapy for suicide prevention (CBTSP): treatment model, feasibility and acceptability. *J Am Acad Child Adolesc Psychiatry* 2009; 48 [10]:1005–13.
- ^b Rotheram-Borus MJ, et al. Enhancing treatment adherence with aspecialized emergency room program for adolescent suicide attempters. *J Am Acad Child Adolesc Psychiatry* 1996; 35:654–663.
- ^c Linehan MM, et al. Cognitive-behavioral treatment of chronically parasuicidal borderline patients. *Arch Gen Psychiatry* 1991; 48:1060–1064.
- ^d Linehan MM. Skills training manual for treating borderline personality disorder. New York: Guilford Press, 1993.
- ^e Miller, AL, et al. Dialectical behavior therapy with suicidal adolescents. New York: Guilford Press, 2007.
- ^f Morgan HG et al. Secondary prevention of non-fatal deliberate self-harm. The green card study. *BJP* 1993; 163: 111–112.
- ^g Excluded secondary publications: Diamond G, et al. Sexual trauma history does not moderate treatment outcome in Attachment-Based Family Therapy (ABFT) for adolescents with suicide ideation. *J Fam Psychol* 2012; 26(4): 595–605; Shpigiel MS, et al. Changes in parenting behaviors, attachment, depressive symptoms, and suicidal ideation in attachment-based family therapy for depressive and suicidal adolescents. *J Marital Fam Ther* 2012; 38(Suppl 1): 271–83.
- ^h Reynolds WM. Suicidal Ideation Questionnaire: Professional Manual. Psychological Assessment Resources Inc., 1987.
- ⁱ Classified as CBT in the meta-analysis.
- ^j Excluded secondary publication: Harrington R, et al. Deliberate self-poisoning in adolescence: why does a brief family intervention work in some cases and not others? *J Adolesc* 2000; 23(1): 13–20.
- ^k Carter GL, et al. Postcards from the EDge project: randomised controlled trial of an intervention using postcards to reduce repetition of hospital treated deliberate self poisoning. *BMJ* 2005; 331: 805–7.
- ^l Henggeler S, et al. Serious Emotional Disturbance in Children and Adolescents: Multisystemic Therapy. New York: Guilford Press, 2002.
- ^m Schmidt U, Davidson KM. Life After Self-Harm: A Guide to the Future. Routledge, 2004.
- ⁿ Hodges K. Child and Adolescent Functional Assessment Scale. Ypsilanti: Eastern Michigan University, 1989.
- ^o Excluded secondary publication: Mehlum L, et al. Dialectical behavior therapy for adolescents with repeated suicidal and self-harming behavior: a randomized trial. *J Am Acad Child Adolesc Psychiatry* 2014; 53(10): 1082–91.
- ^p Shochet I, et al. Resourceful Adolescent Parent Program: group leader's manual. Brisbane, Australia: Griffith University, 1997.
- ^q Excluded secondary publication: Wingate LR, et al. (Comparison of compensation and capitalization models when treating suicidality in young adults. *J Consult Clin Psychol*, 2005. 73(4): 756–62.
- ^r Miller I, et al. (1986). The modified scale for suicidal ideation: Reliability and validity. *J Consult Clin Psychol*, 54, 724–725.
- ^s Excluded secondary publications: Slee N, et al. Emotion regulation as mediator of treatment outcome in therapy for deliberate self-harm. *Clin Psychol Psychother* 2008; 15(4): 205–16.; Spinhoven P, et al. Childhood sexual abuse differentially predicts outcome of cognitive-behavioral therapy for deliberate self-harm. *J Nerv Ment Dis* 2009; 197(6): 455–7.
- ^t Osman A, et al. The Reasons for Living Inventory for Adolescents (RFL-A): development and psychometric properties. *J Clin Psychol* 1998; 54: 1063–1078.

Table 2

Study characteristics: Non-randomized controlled trials conducted in clinical settings (N = 19).

Study; country	Study design; level of evidence	Target population	Participants	Intervention description	Comparison condition	Risk of bias	Suicide related outcome(s) assessed; Longest follow-up	Results	Interpretation
Asarnow et al. (2015) [73] USA	Study design: Pre-test/post-test case series Level of evidence: IV	Inclusion: SA in past 3 months; stable living situation Exclusion: No contact information available for follow-up; psychosis; substance abuse/dependence; not English-speaking; no family to participate Recruited from: Hospital/ED	N = 35 Mean age: 14.89 (SD: 1.6; Range: 11-18) Gender: 14% male	Suicide-specific family-based cognitive behavioral therapy comprising psycho-education plus individual therapy. The Safe Alternatives for Teens & Youths program (SAFETY Program) Length: Up to 20 sessions over 12 weeks, incl: 1 x family session then individual (16 x youth-only & parent-only), then up to 16 x family sessions Developed by: Henggeler (2002) Delivered by: a MH professional	NA	Adequately powered: NR Outcome assessor blinding: NA Less than 15% drop-out rate: Yes (11.4%) Use of statistical testing to measure change from pre-test to post-test: Yes	SI: Harkavy-Asnis Suicide Survey, passive suicidal ideation subscale. SA: Harkavy-Asnis Suicide Survey, suicide attempt subscale. SRB: Harkavy-Asnis Suicide Survey, active suicidal behavior and ideation subscale. Longest follow-up: 6 months post-intervention	SI: Pre-test Mean (SD): 12.69 (9.79) Post-test Mean (SD): 9.19 (10.14) SA: Pre-test Mean (SD): 0.89 (1.86) Post-test Mean (SD): 0.13 (0.34) SRB: Pre-test Mean (SD): 3.71 (4.42) Post-test Mean (SD): 1.81 (2.69)	There was evidence of a significant reduction in SI (t -test = 2.56, p = 0.016, Cohen's d = 0.39), SA (t -test = 2.42, p = 0.019), and SRB (t -test = 2.63, p = 0.013) between baseline and three-month follow-up. Four young people either re-attempted suicide and/or re-engaged in NSSI during the treatment period (significance test not reported).
Brent et al. (2009) [91] USA	Study design: Non-randomized, experimental trial Level of evidence: III-2	Inclusion: Had major unipolar mood disorder & SA in past 90 days; living with a parent or guardian who could participate in treatment Exclusion: Substance dependence, bipolar disorder, psychosis, or developmental disorder Recruited from: Unclear	Whole sample N = 124 Mean age: 15.8 (SD: 1.5; Range: 12-18) Gender: 22.6% male Treatment group (1) N = 18 Age/gender: NR Treatment group (2) N = 93 Age/gender: NR Control group N = NR Age/gender: NR N = 61	Suicide-specific individual cognitive behavioral therapy with some elements of dialectical behavior therapy. Length: between 12 and 16 weekly sessions. Developed by: Study authors Delivered by: Unclear	Medication management or combined medication & CBT	Adequately powered: No Outcome assessor blinding: Yes Less than 15% drop-out rate: No (33.1%) Use of statistical testing to measure change from pre-test to post-test: Yes	SA: Columbia Classification Algorithm of Suicide Assessment SRB: Columbia Classification Algorithm of Suicide Assessment. Suicide: not described. Longest follow-up: Post-intervention only	SA: NR SRB: NR Suicide: NR	There was evidence of an increase in SRB between baseline and six-month follow-up in the combination (i.e., psycho- and pharmacotherapy group) compared to either condition alone (22/93 vs. 2/31; Fisher's exact test p = <0.04). There was one completed suicide after the six-month follow-up, however, it is unclear to which treatment group this young person had been allocated.
Courtney & Flament, (2015) [74] Canada	Study design: Pre-test/post-test case series Level of evidence: IV	Inclusion: BPD features with SI OR SH in past 4 months Exclusion: psychosis; developmental disorder Recruited from: MH outpatient	N = 61 Mean age: 16.5 (SD: 0.8; Range: 15-18) Gender: 7% male	Dialectical behavior therapy adapted for adolescents in tertiary care. A-DBT-A Length: 1 x weekly group-based and 1 x weekly individual sessions over 14-weeks (session duration not stated). Developed by: Based on Miller et al. (2006) but adapted by the study authors Delivered by: a MH professional	NA	Adequately powered: NR Outcome assessor blinding: NA Less than 15% drop-out rate: No (49.2%) Use of statistical testing to measure change from pre-test to post-test: Yes	SI: Suicidal Ideation Questionnaire (SIQ). SRB: Medical/clinical records. Suicide: NR Longest follow-up: Post-intervention only	SI: Pre-test Median (IQR): 131.0 (92.0 to 144.0). Post-test Median (IQR): 77.0 (48.5 to 121.0). SRB: NR Suicide: NR	There was evidence of a significant reduction in SI (t -test = 4.96, p < 0.001, Cohen's d = 0.89) between baseline and the 15-week post-intervention assessment. There was also evidence of a significant reduction in the proportion of young people engaging in SRB over this period (36/42 vs. 16/42, McNemar test p < 0.001). There were no reports of completed suicides.
Cwik et al. (2016) [82] USA	Study design: Pre-test/post-test case series Level of evidence: IV	Inclusion: Apaches with SA in past 90 days Exclusion: none Recruited from: Community suicide surveillance system	N = 13 Mean age: 14.3 (SD: 2.2) Gender: 8% male	New Hope, a brief psycho-education intervention for American Indian adolescents Length: 1-2 visits (2-4 h total). Developed by: Study authors Delivered by: Community Mental Health Workers	NA	Adequately powered: No Outcome assessor blinding: NA Less than 15% drop-out rate: No (15.4%) Use of statistical testing to measure change from pre-test to post-test: Yes	SI: SIQ Longest follow-up: 3 months Post-intervention	SI: N (%) scoring above clinical cut-off: Pre-test: 7/11 (64%) Post-test: 1/10 (10%)	The number of participants who scored above the clinical cut-off for the SIQ seemed to decrease over the follow-up period.

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Table 2 (continued)

Study; country	Study design; level of evidence	Target population	Participants	Intervention description	Comparison condition	Risk of bias	Suicide related outcome(s) assessed; Longest follow-up	Results	Interpretation
Diamond et al. (2013) [83] ^a USA	Study design: Pre-test/post-test case series Level of evidence: IV	Inclusion: LGB discharged from hospital with SI (admitted for SI or SA) Exclusion: Psychosis or ID Recruited from: Hospital/ED	N = 10 Mean age: 15.1 (SD: 1.37; Range: 14–18) Gender: 20% male	Attachment-based family therapy adapted for use with suicidal LGB youth. <i>ABFT-LGB</i> Length: 12 x weekly sessions (range = 8–16). Sessions lasted for 60-min & sessions 3–5 were for parents only. Developed by: Study authors Delivered by: a MH professional	NA	Adequately powered: No Outcome assessor blinding: NA Less than 15% drop-out rate: Yes (0.0%) Use of statistical testing to measure change from pre-test to post-test: Yes	SI: SIQ-Junior (SIQ-JR) Longest follow-up: Post-intervention only	SI: Pre-test Mean (SD): 51.00 (13.00) Post-test Mean (SD): 6.88 (7.34)	There was evidence of a significant reduction in SI between baseline and the 3-month post-intervention assessment (F-test = 18.78, $p =$ 0.001, Cohen's $d = 0.21$).
Duarte-Velez et al. (2016) [75] Puerto Rico	Study design: Pre-test/post-test case series Level of evidence: IV	Inclusion: Admitted to ED with SI or SA, hospitalized, stabilized and referred to outpatient; legal guardian. Exclusion: Psychosis; developmental disorder; ID; already receiving psychotherapy; involvement in a legal procedure that would require psychological care mandated by the judicial system Recruited from: Hospital/ED	N = 11 Mean age: 15.36 (SD-NR; Range: 13–17) Gender: 45% male	Cognitive behavioral therapy adapted for Puerto Rican adolescents with suicidal behavior. Length: Weekly individual sessions lasting for 1 h & delivered over 6 months. Plus 60–120 min family sessions & follow-up bi-weekly as necessary. Phone calls & case management as needed. Developed by: Study authors Delivered by: a MH professional	NA	Adequately powered: No Outcome assessor blinding: NA Less than 15% drop-out rate: No (27.3%) Use of statistical testing to measure change from pre-test to post-test: Yes	SI: SIQ-JR Longest follow-up: Post-intervention	SI: Pre-test Mean (SD): 27.20 (NR) Post-test Mean (SD): 16.00 (NR)	There was evidence of a reduction in SI between baseline and the six month post-intervention assessment (significance test not reported).
Esposito- Smythers et al. (2006) [76] USA	Study design: Pre-test/post-test case series Level of evidence: IV	Inclusion: Admitted to inpatient unit for SI/SA with co-occurring alcohol abuse/dependence Exclusion: ID, DSM-IV dependence on substances other than alcohol or cannabis. Recruited from: Hospital/ED	N = 6 Mean age: 15 (SD: 1; Range: 14–16) Gender: 17% male	Integrated cognitive behavioral therapy for adolescents with co-occurring alcohol use disorder and suicidality. Length: Acute phase: Weekly sessions lasting 1 h & delivered over 6 months (plus maintenance & booster phases). Developed by: Study authors, incorporating modifications of Monti's (2002) ^b coping skills training package for youth with co-occurring alcohol use disorder Delivered by: a MH professional	NA	Adequately powered: No Outcome assessor blinding: NA Less than 15% drop-out rate: No (16.7%) Use of statistical testing to measure change from pre-test to post-test: No	SI: SIQ SRB: NR Longest follow-up: 12 months post-intervention	SI: Pre-test Mean (SD): 80.80 (NR) Post-test Mean (SD): 32.80 (NR)	There was evidence of a reduction in SI between baseline and the 12 month post-intervention assessment (significance test not reported). Two young people re-engaged in SRB during this period (significance test not reported).
Geddes et al. (2013) [77] Australia	Study design: Pre-test/post-test case series Level of evidence: IV	Inclusion: At least 3 BPD features & SI/SH in past 12 months Exclusion: Primary diagnosis of psychosis or substance abuse; ID Recruited from: MH Outpatient	N = 6 Mean age: 15.1 (SD-NR; Range 14–15) Gender: 0% male	Dialectical behavior therapy modified for adolescents: Life Surfing Length: 1–2 weekly sessions lasting for 1 h & delivered over 26 weeks. Plus a weekly 2 h family skills group delivered over an 18-week period. Developed by: Based on Swales (2000) ^c but adapted by the study authors. Delivered by: NR	NA	Adequately powered: No Outcome assessor blinding: NA Less than 15% drop-out rate: No (16.7% by the three-month follow-up period) Use of statistical testing to measure change from pre-test to post-test: Yes	SI: NR SBR: Self-Harm/Suicidal Thoughts Questionnaire: Parent and Adolescent Versions. SA: NR Longest follow-up: 12 months post-baseline	SA: NR SRB: NR	There was evidence of a reduction in the proportion of young people reporting SI between baseline and the 18-week post-intervention assessment (significance test not reported). By the 18-week post-intervention assessment, 5 of the 6 young people had had no further episodes of SRB, whilst the sixth reported a 50% reduction in SRB frequency (significance tests not provided). By the 12 month follow-up

Gutstein & Rudd (1990) [78] USA	Study design: Pre-test/post-test case series Level of evidence: IV	Inclusion: Referred to a guidance center following a near-lethal SA/persistent suicide threats (severe risk) Exclusion: NA Recruited from: Hospital/Ed, MH outpatient & community	N = 47 Mean age: 14.4 (SD-NR; Range: 7-19) Gender: 47% male	A suicide-specific intensive group crisis intervention: Systemic Crisis Intervention Program Length: Two × 4-hour group meetings over a 2-6 week period. Developed by: Study authors Delivered by: NR	NA	Adequately powered: No Outcome assessor blinding: NA Less than 15% drop-out rate: Yes (0.0%) Use of statistical testing to measure change from pre-test to post-test: No	SA: Parental report Longest follow-up: Post-intervention only	SA: NR	period, no young person had a further episode of SA (significance test not reported). There was evidence of a reduction in the proportion of young people engaging in SA between baseline and the 18 month follow-up assessment (significance test not reported).
James et al. (2011) [79] UK	Study design: Pre-test/post-test case series Level of evidence: IV	Inclusion: Living in 'out of home care' & engaged in SH for >6 months Exclusion: diagnosis of schizophrenia, bipolar disorder, autism spectrum disorder; Moderate-severe mental impairment Recruited from: MH outpatient service	N = 25 Mean age: 15.5 SD: 1.5; Range: 13-17 Gender: 12% male	Dialectical behavior therapy comprising a skills training group, individual therapy, telephone support, support for schools/carers & outreach. Length: 1-hour individual sessions plus 2-hour group sessions delivered weekly over 12 months. Developed by: Based on Linehan (1993) and Rathus and Miller (2002) ^d but adapted by the study authors. Delivered by: a MH professional	NA	Adequately powered: No Outcome assessor blinding: NA Less than 15% drop-out rate: No (28.0%) Use of statistical testing to measure change from pre-test to post-test: Yes	SRB: Clinical interview Longest follow-up: Post-intervention only	SRB: NR	There was evidence of a reduction in the proportion of young people engaging in SRB between baseline and the 12 week post-intervention period (14/18 young people had ceased engaging in SRB altogether) (significance tests not provided). There was also evidence of a reduction in the frequency of these SRB episodes over this period (significance tests not provided).
James et al. (2015) [80] UK	Study design: Pre-test/post-test case series Level of evidence: IV	Inclusion: SH in past 12 months Exclusion: NR Recruited from: MH outpatient & community	N = 154 Mean age: 14.9 (SD: 1.3; Range: 12-18) Gender: 14.8% male	Dialectical behavior therapy for adolescents Length: Three-hour group sessions delivered twice weekly, plus weekly individual or family sessions, 30-60 min in duration. Delivered over 16 weeks. Developed by: Based on Miller (2006) Delivered by: MH professional	NA	Adequately powered: NR Outcome assessor blinding: NA Less than 15% drop-out rate: No (30.3%) Use of statistical testing to measure change from pre-test to post-test: Yes	SRB: Youth Outcome Questionnaire, Self-Report, version 2.0, item 21 Longest follow-up: Post-intervention only	SRB: Pre-test Mean (SD): 2.06 (1.68). Post-test Mean (SD): 0.65 (0.98).	There was evidence of a significant reduction in SRB between baseline and the 16-week post-intervention assessment (F-test = 68.83, $p <$ 0.001, $\eta^2 = 0.42$).
Katz et al. (2004) [84] Canada	Study design: Non-randomized, experimental trial Level of evidence: III-2	Inclusion: Admitted to Inpatient unit for SA or SI Exclusion: ID, severe learning disability, psychosis, bipolar disorder Recruited from: Hospital/ED	Whole sample N = 62 Mean age: 15.4 (Range: 14-17) Gender: 16.1% male Treatment group N = 31 Age/gender: NR Control group N = 31 Age/gender: NR	Individual & group dialectical behavior therapy Length: 10 daily group sessions plus 4 individual sessions delivered over 2 weeks Developed by: Based on Miller (1997) ^e but adapted by the study authors Delivered by: MH professional	TAU: daily psychody- namic psychother- apy group, weekly individual therapy, and psychodyna- mically-- oriented milieu.	Adequately powered: Study authors provide power calculations, however, study unlikely to be adequately powered for SRB. Outcome assessor blinding: NA Less than 15% drop-out rate: Yes (10.0%) Use of statistical testing to measure change from pre-test to post-test: Yes	SI: SIQ-JR Suicide: NR Longest follow-up: 1 year post-intervention	SI: Post-intervention Intervention mean (SD): 40.90 (24.73) Control mean (SD): 37.97 (24.56) 12 months: Intervention mean (SD): 18.15 (12.52) Control mean (SD): 19.25 (17.89) Suicide: NR	There was no evidence of a reduction in SI between the intervention and control groups at post-intervention (40.90 ± 24.73 vs. 37.97 ± 24.56) and at the 12 month follow-up assessment (18.15 ± 12.52 vs. 19.35 ± 17.89). There were no completed suicides in either group by the 12 month follow-up assessment (significance test not reported).
King et al. (2003) [81] Australia	Study design: Pre-test/post-test case series Level of evidence: IV	Inclusion: Called helpline and reported SI Exclusion: None Recruited from: Telephone helpline	N = 101 Age: NR Gender: Unclear	Kids helpline Single crisis phone call Length: Mean duration 40 min; range 10-120 min Developed by: Charitable organization Delivered by: trained volunteers	NA	Adequately powered: NR Outcome assessor blinding: NA Less than 15% drop-out rate: Yes (0.0%)	SI: Idiosyncratic, binary-coded instrument adapted from items from the Mini International Neuropsychiatric Interview.	SI: Pre-test Mean (SD): 6.30 (2.22). Post-test Mean (SD): 3.01 (2.43).	There was evidence of a significant reduction in SI from the beginning to the end of the call (average call duration 40 min) (t -test = 12.66, $p <$ 0.005, $\eta^2 = 0.62$).

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Table 2 (continued)

Study; country	Study design; level of evidence	Target population	Participants	Intervention description	Comparison condition	Risk of bias	Suicide related outcome(s) assessed; Longest follow-up	Results	Interpretation
Law et al. (2016) [85] Hong Kong	Study design: Non-randomized, experimental trial Level of evidence: III-2	Inclusion: Admitted to the ED with SH Exclusion: any DSM IV-TR Axis II disorder; psychosis; bipolar disorder Recruited from: Hospital/ED	Whole sample N = 78 Mean age: NR Range: 18–34 Gender: NR Treatment group N = 40 Mean age: 24.7 (SD: 5.4) Gender: 18.4% male Control group N = 38 Mean age: 26.0 (SD: 6.2) Gender: 11.1% male	Brief contact intervention: Volunteer mentorship Length: ≥2 contacts per month over 9 months. Developed by: Study authors Delivered by: trained volunteers supervised by psychiatrists, psychologists, and social workers	TAU (not described)	Use of statistical testing to measure change from pre-test to post-test: Yes Adequately powered: Study authors provide power calculations, however, study unlikely to be adequately powered for SRB Outcome assessor blinding: NA Less than 15% drop-out rate: No (67.6%) Use of statistical testing to measure change from pre-test to post-test: Yes Adequately powered: No Outcome assessor blinding: NA Less than 15% drop-out rate: Yes (0.0%) Use of statistical testing to measure change from pre-test to post-test: Yes	Longest follow-up: Post-intervention only SI: SIQ SRB: Hospital records Longest follow-up: Post-intervention only	SI: Post-intervention: Intervention mean (SD): 20.70 (3.00) Control mean (SD): 15.60 (6.50) SRB: NR	There was no evidence of a significant reduction in SI between the intervention and control groups at post-intervention (20.70 ± 3.00 vs. 15.60 ± 6.50, $\beta = 2.31$, SE = 2.52, $p > 0.05$). There was also no evidence of a reduction in SRB between the intervention and control groups by this time point (4/38 vs. 4/36) (significance test not provided).
Oldershaw et al. (2012) [86] UK	Study design: Retrospective cohort study Level of evidence: III-2	Inclusion: Reported history of SH Exclusion: ID; serious head injury; used sedatory side effects; primary diagnosis not depression or SH. Recruited from: MH outpatient, schools & personal contacts	Whole sample N = 33 Mean age: NR (SD: NR; Range: 12–18) Gender: NR Treatment group N = 24 Age: NR Gender: 4.2% male Control N = 9 Age: NR Gender: 22.2% male	Standalone, formulation based, and modularized cognitive behavioral therapy with core and optional modules, depending on clinical need. Length: 12 sessions Developed by: Study authors Delivered by: MH professional	No treatment: Participants either declined or did not pursue treatment	Adequately powered: No Outcome assessor blinding: NA Less than 15% drop-out rate: Yes (0.0%) Use of statistical testing to measure change from pre-test to post-test: Yes	SRB: Idiosyncratic, binary-coded instrument Longest follow-up: 5 months post-baseline	SRB: NR	There was evidence of a significant reduction in the proportion of participants engaging in SRB between the intervention and control groups at post-intervention (14/24 vs. 3/9). There was also evidence of a significant reduction in the frequency of SRB by this time point ($Z = -3.20$, $p < 0.001$).
Perera Ramani & Kathirarachchi, (2011) [87] Sri Lanka	Study design: Non-randomized, experimental trial Level of evidence: III-2	Inclusion: Admitted to hospital for SA; categorized as medium- and low-intent Exclusion: Diagnosed with major psychiatric disorder Recruited from: Hospital/ED	Whole sample N = 124 Mean age: NR (SD: NR; Range: 15–24) Gender: Unclear Treatment group N = 62 Age/Gender: NR Control group N = 62 Age/Gender: NR	Individual problem solving therapy Length: 4 sessions delivered over 1 month Developed by: Based on Palmer (1995) ^f Delivered by: MH professional	TAU: routine care (referral to a medical officer, psychiatric referral, referrals to other agencies).	Adequately powered: No Outcome assessor blinding: NR Less than 15% drop-out rate: No (18.5%) Use of statistical testing to measure change from pre-test to post-test: No	SA: NR Longest follow-up: 6 months post-baseline	SA: NR	There was a reduction in the proportion of participants engaging in SA between the intervention and control groups at post-intervention (0/55 vs. 2/46) (significance test not reported).
Rathus & Miller, (2002) [88] USA	Study design: Non-randomized, experimental trial Level of evidence: III-2	Inclusion: SA or SI in past 4 months AND Borderline Personality Disorder features Exclusion: NR Recruited from: MH outpatient	Whole sample N = 111 Age: NR Gender: 21.6% male Treatment group N = 29 Mean age: 16.1 (SD: 1.2; Range: NR) Gender: 7% male Control group N = 82 Mean age: 15.0 (SD:	Dialectical behavior therapy adapted for adolescents. Length: Two sessions per week for 12 weeks Developed by: Based on Linehan (1993) but adapted for adolescents by study authors Delivered by: MH professional	Active placebo: Short term psychodynamic or supportive approach aimed at resolving acute problems.	Adequately powered: No Outcome assessor blinding: NA Less than 15% drop-out rate: Unclear Use of statistical testing to measure change from pre-test to post-test: Yes	SI: Beck Scale for Suicidal Ideation (BSSI) SA: Self-report Longest follow-up: 3 months post-intervention	SI: Pre-test Mean (SD): 9.80 (5.30) Post-test Mean (SD): 3.80 (4.60) SA: NR	There was a significant reduction in SI between baseline and the 12-week post-intervention assessment (t -test = 2.65, $p = 0.026$). There was also evidence of a reduction in SA between the intervention and control groups by the 12-week post-intervention assessment (1/29 vs. 7/82).

Rotheram-Borus et al. (1996) [89]; (2000) [35] USA	Study design: Historical controlled study Level of evidence: III-3	Inclusion: Presented to ED with SA & hospitalized for <1 week Exclusion: Low IQ, no parent or family Recruited from: Hospital/ED	1.7; Range: NR) Gender: 27% male Whole sample N = 140 Mean age: 15.0 (SD: NR; Range: 12-18) Gender: 0% male Treatment group N = 65 Mean age: 15.0 (SD: 1.4) Control group N = 75 Mean age: 15.3 (SD: 1.6)	Specialized Emergency Room Program: Comprised 1 family psychotherapy session plus psycho-education video. Length: Session = NR; video = 20 min Developed by: Study authors Delivered by: MH professional	TAU: evaluation to determine if hospitalization required & referral to outpatient therapy.	Adequately powered: Likely to be adequately powered for SI. Outcome assessor blinding: NA Less than 15% drop-out rate: Yes (0.0%) Use of statistical testing to measure change from pre-test to post-test: Yes	SI: Harkavy-Asnis Suicide Survey, passive suicidal ideation subscale SA: Self- and parental-report in conjunction with hospital records Longest follow-up: 18 months post-suicide attempt	SI: Post-intervention: Intervention mean (SD): 1.40 (2.38) Control mean (SD): 2.10 (2.86) SA: 18-month follow-up: Tx - 6/65 participants reattempted; Control - 11/75 participants reattempted	There was evidence of a significant reduction in SI between the intervention and control groups by the post-intervention assessment ($p < 0.001$). There was also evidence of a reduction in SA between the intervention and control groups by the 18 month follow-up assessment (significance test not reported).
Wharff et al. (2012) [90] USA	Study design: Historical controlled study Level of evidence: III-3	Inclusion: Presented to the ED with SRB Exclusion: not living with family; presented to ED without a family member; intoxicated/sedated at time of presentation; psychosis or developmental delay; presented during overnight shift or on weekend Recruited from: Hospital/ED	Whole sample N = 250 Mean age: NR (SD: NR; Range: 13-18) Gender: 29% male Treatment group N = 100 Mean age: 15.6 (SD: 1.5) Gender: 24% male Control group N = 150 Age: NR Gender: 26% male	One session of family based crisis intervention Developed by: Study authors Length: NR Delivered by: MH professional	TAU: retrospective comparison group who presented to the same ER prior to implementation of FBCL.	Adequately powered: No Outcome assessor blinding: Unclear Less than 15% drop-out rate: No (44.6%) Use of statistical testing to measure change from pre-test to post-test: No	SRB: NR SA: NR Suicide: NR Longest follow-up: 3 months post-intervention	SRB: NR SA: NR Suicide: NR	There were no episodes of SA or completed suicide in either the intervention or control group by the three month follow-up period. Two participants (unclear if allocated to intervention or control groups) were hospitalized for SRB by the three month follow-up assessment.

Notes: ED = Emergency Department; ID = Intellectual Disability; ITT = intention-to-treat; IQR = Interquartile Range; MH = mental health; NR = not reported; TAU = treatment as usual; SA = suicide attempt; SD = standard deviation; SH = self-harm; SI = suicidal ideation; SRB = suicide-related behavior.

^a Corrected version of the same paper published in 2012.

^b Monti PM, et al. Treating Alcohol Dependence: A Coping Skills Training Guide. 2nd ed. New York, NY: Guilford Press; 2002.

^c Swales M, et al. Linehan's Dialectical Behaviour Therapy (DBT) for borderline personality disorder: overview and adaptation. *J Ment Health* 2000; 9(1): 7-23.

^d Rathus J, Miller A. Dialectic behavior therapy adapted for suicidal adolescents. *Suicide Life Threat Behav* 2002; 32: 146-157.

^e Miller AL, et al. Dialectical behavior therapy adapted for suicidal adolescents. *J Pract Psychiatry Behav Health* 1997; 3:78-86.

^f Palmer S, Dryden W. Counseling for Stress Problems. New Delhi: Sage Publications, 1995.

Table 3

Study characteristics: Randomized controlled trials conducted in educational or workplace settings (N = 15).

Study; country	Target population	Participants	Intervention description	Comparison condition	Risk of bias	Suicide related outcome(s) assessed; longest follow-up
Universal interventions Guille et al. (2015) [113] USA	Inclusion: Medical students beginning their internship in July 2009 or July 2011 at one of two participating university hospitals Exclusion: NR Recruited from: Universities (N = 2)	Whole sample N = 199 Mean age: 25.2 (SD: 8.1; Range: NR) Gender: 50.7% male Treatment group N = 100 Mean age: 24.9 (SD: 8.7) Gender: 49% male Control group N = 99 Mean age: 25.4 (SD: 7.4) Gender: 51.6% male	Individual access to MoodGym: online Cognitive Behavioral Therapy. Module 1: Understanding the interplay between thoughts, emotions & behavior. Modules 2–3: Cognitive restructuring. Module 4: Problem-solving. Length: Four 30-min modules Developed by: National Institute for Mental Health Research at The Australian National University Delivered by: Self-directed	Active placebo: Email once a week for 4 weeks with information about depression, suicide & where to seek treatment.	Random sequence generation method: NR Allocation concealment method: Independent researcher Ascertainment of SH repetition: Self-report Outcome assessor blinding: NA Less than 15% drop-out rate: Yes (0.0%) Was ITT analysis undertaken: Yes	SI (dichotomous): Item 9 of the Patient Health Questionnaire-9 (PHQ-9) Longest follow-up: 12 months post-intervention
Orbach & Bar-Joseph, (1993) [92] Israel	Inclusion: High school juniors from six schools Exclusion: None Recruited from: Secondary schools (N = 6)	Whole sample N = 393 Mean age: NR Gender: 45% male Treatment group N = 215 Age/gender: NR Control group N = 178 Age/gender: NR	Weekly group psycho-education workshops: 1) Depression & happiness 2) The individual & their family 3) Helplessness 4) Coping with failure 5) Coping & problem solving 6) Coping with suicidal urges 7) Summary Length: Seven 2-hour workshops Developed by: based on Ross (1997) ^a and adapted by study authors. Delivered by: NR	TAU: social issues discussion class	Random sequence generation method: NR Allocation concealment method: NR Ascertainment of SH repetition: Self-report Outcome assessor blinding: NA Less than 15% drop-out rate: Yes (0.0%) Was ITT analysis undertaken: Mixed methods	SI (continuous): Israeli Index of Potential Suicide (IIPS) Longest follow-up: Post-intervention only
Till et al. (2017) [119] Austria	Inclusion: medical, psychology and communication studies undergraduate students Exclusion: None Recruited from: University (N = 1)	Whole sample N = 161 Mean age: 24.5 (SD 5.8) Gender: 32.9% male Treatment group N = 121 Mean age: 24.3 (SD NR) Gender: 33.9% male Control group N = 40 Mean age: 25.0 (SD 6.8) Gender: 30% male	Psychoeducation plus support Three German-language websites on suicide-related education and prevention. Two of the three websites also offered email counseling by peers. Length: NA Developed by: mental health organizations Delivered by: Self-directed	A website unrelated to suicide or mental health	Random sequence generation method: NR Allocation concealment method: NR Ascertainment of DSH repetition: NA Outcome assessor blinding: NA Less than 15% drop-out rate: Unclear Was ITT analysis undertaken: Yes	SI (continuous): Reasons for Living Inventory (RFLI) Longest follow-up: Post-intervention only
Indicated interventions Eggert et al. (2002) ^b [95], USA	Inclusion: Students who screened positive for SRB Exclusion: None Recruited from: Secondary schools (N = 7)	Whole sample N = 341 Mean age (SD): NR (Range: 14–19) Gender: 48% male Treatment group 1 (C-CAST) N = 103 Mean age: 16.02 (SD: 1.14) Gender: 40.77% male Treatment group 2 (C-Care) N = 117 Mean age: 15.71 (SD: 1.21)	Supportive intervention comprising: 1) C-CARE: One individual assessment interview followed by one counseling session & social connections intervention with parents and school staff	TAU: a brief assessment interview and social connections intervention with parents and school personnel.	Random sequence generation method: Block randomization using a predetermined sequence Allocation concealment method: NR Ascertainment of SH repetition: Self-report Outcome assessor blinding: NA Less than 15% drop-out: No (20.5%)	SRB: High School Questionnaire: Profile of Experiences (HSQ) Longest follow-up: 9 months post-baseline Not included in MA

		Gender: 52.14% male Control group N = 121 Mean age: 15.62 (SD: 1.26) Gender: 50.83 male	2) C-CARE plus a small group prevention program.Length:		Was ITT analysis undertaken: No	
			1) 2-hour assessment plus one 1.5–2 h counseling; 2) Additional 12 × 1 hour sessions over 6 weeksDeveloped by: study authors Delivered by: 1) Trained research staff e.g. practice nurses & social workers; 2) Teachers, counselors or nurses A one-off suicide-specific problem solving intervention. Included a video, narrated Power-Point presentation & a case study. Included identifying problems and cognitive, behavioral & affective reactions. It encouraged participants to elicit their problems and apply the skills learned in the video to their personal problems. Length: 40 mins Developed by: based on D'Zurilla and Nezu (1999) ^c Delivered by: NR Online cognitive behavioral therapy (Reframe IT) Length: Eight self-directed modules over 10 weeks Developed by: study authors Delivered by: self-directed, in the presence of school well-being staff	Active placebo: Video about health issues e.g. diet, exercise, and sleep.	Random sequence generation method: NR Allocation concealment method: NR Ascertainment of SH repetition: Self-report Outcome assessor blinding: NA Less than 15% drop-out rate: No (31.8%) Was ITT analysis undertaken: No	SI (continuous): Beck Scale for Suicidal Ideation (BSSI) Longest follow-up: 1 month post-baseline
Fitzpatrick et al. (2005) [115] USA	Inclusion: Students who screened positive for SI Exclusion: students who were judged to represent an immediate threat of danger to themselves or others Recruited from: University (N = 1)	Whole sample N = 110 Mean age: 19.02 (SD: 1.21; Range: 18–24) Gender: 45% male Treatment group: NR Control group: NR				
Hetrick et al. (2017) [123] Australia	Inclusion: Presented to school counselor with SI Exclusion: Intellectual disability; psychotic symptoms; inability to speak English Recruited from: Secondary schools (N = 18)	Whole sample N = 50 Mean age: 14.7 (SD: 1.4) Gender: 18% male Treatment group N = 26 Mean age: 14.8 (SD: 1.6) Gender: 19.3% male Control group N = 24 Mean age: 14.5 (SD: 1.3) Gender: 16.7% male		TAU: contact with school wellbeing staff plus any outside mental health service provision normally available.	Random sequence generation method: Online randomization program, stratified by school Allocation concealment method: The online program did not allow knowledge of treatment next to be allocated before the participant details were entered into the computer Ascertainment of SH repetition: Interview Outcome assessor blinding: NA Less than 15% drop-out: No (28.6%) Was ITT analysis undertaken: Yes	SI (continuous): SIQ SA: a specifically designed questionnaire that asked participant whether they had attempted suicide since their last assessment, and if so, how many times Longest follow-up: 3 months post-intervention
Hill & Pettit, (2016) [122] ^d USA	Inclusion: Endorsed a perceived burdensomeness score of 17 or greater on the Interpersonal Needs Questionnaire Perceived Burdensomeness subscale (Van Orden et al., 2012) ^e Exclusion: Current psychosocial treatment or use of psychoactive	Whole sample N = 80 Mean age: 16.9 (SD: 1.7; Range: 13–19) Gender: 31.2% male Treatment group N = 40 Age/gender: NR Control group N = 40	Online cognitive behavioral therapy (LEAP: Learn, Explore, Assess you options, Plan) Length: Two modules delivered over two weeks Developed by: study authors Delivered by: self-directed	Placebo: e-mail containing psychoeducational information about mental health & suicide, and resources for mental health treatment and suicide/crisis counseling.	Random sequence generation method: Sequentially numbered envelopes Allocation concealment method: Sealed, opaque envelopes Ascertainment of SH repetition: Self-report questionnaire.	SI (continuous): BSSI Longest follow-up: 2 months post-baseline

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Table 3 (continued)

Study; country	Target population	Participants	Intervention description	Comparison condition	Risk of bias	Suicide related outcome(s) assessed; longest follow-up
Hooven et al. (2012) [96] USA	medications, unless on a stable dose for >8 weeks Recruited from: Schools (N = NR) & public gathering places Inclusion: Students who met criteria for suicide risk status Exclusion: None Recruited from: Secondary schools (N = 20)	Age/gender: NR Whole sample N = 615 Mean age: 16.0 (SD: NR; Range: 14–19) Gender: 40% male Treatment group C-CARE N = 153 Age/gender: NR P-CARE N = 155 Age/gender: NR Combined N = 164 Age/gender: NR Control group N = 143 Age/gender: NR	Combined intervention comprising: 1) Counselors Care, Assess, Respond and Empower (C-CARE) 2) Parents Care, Assess, Respond and Empower (P-CARE) 3) Combined C-CARE and P-CARE Length: two 2-hour sessions over a 1-month period Developed by: study authors Delivered by: unclear	Placebo: brief screening interview.	Outcome assessor blinding: NA Less than 15% drop-out: Yes (13.8%) Was ITT analysis undertaken: Yes Random sequence generation method: NR Allocation concealment method: NR Ascertainment of SH repetition: Self-report Outcome assessor blinding: NA Less than 15% drop-out rate at post-intervention: Unclear Was ITT analysis undertaken: Unclear	SRB: High School Questionnaire: Profile of Experiences (HSQ) Longest follow-up: 15 months post-baseline Not included in MA
Kovac & Range, (2002) [114] USA	Inclusion: Students who screened positive for SRB Exclusion: None Recruited from: University (N = 1)	Whole sample N = 121 Mean age: 23.12 (SD: 5.44; Range: 18–42) Gender: 27.3% male Treatment group N = NR Age/gender: NR Control group N = NR Age/gender: NR	A writing intervention ^f to examine whether writing with 'cognitive change' reduced suicide risk when compared to writing just about suicidal experience and compared to controls. Group 1: Wrote about being suicidal & were instructed to think about their thoughts and feelings at the time. Group 2: Wrote about being suicidal but were asked to provide details about the event. Group 3: Control. Length: four 20-min sessions delivered once a day for 4 days Developed by: study authors Delivered by: Unclear	Placebo: Wrote in detail about their bedroom	Random sequence generation method: NR Allocation concealment method: NR Ascertainment of SH repetition: Self-report Outcome assessor blinding: NA Less than 15% drop-out rate: No (19.1%) Was ITT analysis undertaken: No	SI (continuous): SIQ Longest follow-up: 6 weeks post-intervention
Pistorello et al. (2012) [116] USA	Inclusion: Students seeking treatment from a University mental health service for SI, SA, or NSSI Exclusion: psychosis, need for inpatient care, or prior DBT treatment Recruited from: University (N = 1)	Whole sample N = 63 Mean age: 20.9 (SD: 1.92) Gender: 19% male Treatment group N = 31 Mean age: 20.4 (SD: 1.6) Gender: 22.6% male Control group N = 32 Mean age: 21.3 (SD: 2.1) Gender: 15.6% male	A combination of individual and group dialectical behavioral therapy. Delivered by: Length: Comprised one 50-min individual psychotherapy session plus a 90-min group skills training session per week, over a 12-month period. Developed by: based on Linehan, (1993) ^g Delivered by: MH professionals	Enhanced TAU: included weekly individual & group therapy, weekly group supervision for therapists, & between-session consultation and family. Interventions as needed.	Random sequence generation method: Computer generated adaptive randomization Allocation concealment method: NR Ascertainment of SH repetition: Interview Outcome assessor blinding: NA Less than 15% drop-out rate at post-intervention: No (22.2%) Was ITT analysis undertaken: Yes. All participants with	SI (continuous): Suicidal Behaviors Questionnaire (SBQ-23) SA: SBQ-32 Longest follow-up: 18 months post-baseline

Robinson W et al. (2016) [97] USA	Inclusion: Students who screened positive for SRB Exclusion: None Recruited from: Secondary schools (N = 4)	Whole sample N = 330 Mean age: NR Range: 14–17 + Gender: 40% male Treatment group: NR Control group: NR	Group coping with stress course: 1) Identifying feelings of stress 2) Reducing negative cognitions & increasing positive thoughts 3) Identifying risk factors for stress 4) Enhancing competencies for managing stress 5) Planning for stress. Length: Fifteen 45-min sessions Developed by: based on Robinson & Case (2003) ^h and adapted by study authors Delivered by: MH professional	TAU: one-to-one sessions on stress management.	missing data were coded as unimproved. Random sequence generation method: NR Allocation concealment method: NR Ascertainment of SH repetition: Self-report Outcome assessor blinding: NA Less than 15% drop-out rate: Yes (0.6%) Was ITT analysis undertaken: NR	SI (categorical): Four suicidality screening items (not included in MA) Longest follow-up: Post-intervention only Not included in MA
Tang et al. (2009) [98] Taiwan	Inclusion: Students with moderate–severe depression, SI, SA, moderate–severe anxiety, or significant hopelessness in previous 2 weeks. Exclusion: acute psychotic symptoms, act out lethal suicidal behaviors, lack proper care for suicide risk by their family, drug abuse, or serious medication condition Recruited from: Secondary schools (N = 1)	Whole sample N = 73 Mean age: NR (Range: 14–18) Treatment group N = 35 Mean age: 15.26 (SD: 1.7) Gender: 34% male Control group N = 38 Mean age: 15.24 (SD: 1.65) Gender: 34% male	Interpersonal psychotherapy Length: Two sessions per week for 6 weeks Developed by: based on Mufson et al. (2004) ⁱ and adapted by study authors Delivered by: School counselor & intern counseling psychotherapists	TAU: psycho-education and individual supportive counseling once or twice a week.	Random sequence generation method: NR Allocation concealment method: NR Ascertainment of SH repetition: Self-report Outcome assessor blinding: NA Less than 15% drop-out rate: Yes (0.0%) Was ITT analysis undertaken: Yes	SI (continuous): BSSI Longest follow-up: Post-intervention only
Multi-modal interventions Schilling et al. (2014) [100] USA	Inclusion: Middle school students Exclusion: None Recruited from: Middle schools (N = 8)	Whole sample N = 470 Age: NR Gender: 47.4% male Treatment group: NR Control group: NR	Suicide-specific psycho-education (plus screening). Signs of Suicide (SOS). Video & discussion guide depicting signs of suicidality & depression and recommended ways to respond. Also included screening to identify students at risk. Length: Video = 17 min Developed by: Screening for Mental Health Inc. Delivered by: Teachers	Attended class as usual & received the program after the study period	Random sequence generation method: Cluster simple randomization. However, 2 schools allocated to control crossed over into the intervention Allocation concealment method: NR Ascertainment of SH repetition: Self-report Outcome assessor blinding: NA Less than 15% drop-out rate: Unclear Was ITT analysis undertaken: No	SRB: Centers for Disease Control and Prevention's (CDC) Youth Risk Behavior Survey (YRBS) Longest follow-up: 3-months post-intervention Not included in MA
Schilling et al. (2016) [99] USA	Inclusion: ninth-grade students Exclusion: None Recruited from: Secondary schools (N = 16)	Whole sample N = 1272 Mean age: NR (Range 14–15) Gender: 58.3% male Treatment group N = 719 Age: NR Gender: 55.8% Control N = 553	Suicide-specific psycho-education (plus screening). Signs of Suicide (SOS). As above.	Waitlist control	No Random sequence generation method: Cluster simple randomization. Allocation concealment method: NR Ascertainment of SH repetition: Self-report Outcome assessor blinding: NA Less than 15% drop-out rate:	SI (dichotomous): CDC YRBS Longest follow-up: Post-intervention only

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Table 3 (continued)

Study; country	Target population	Participants	Intervention description	Comparison condition	Risk of bias	Suicide related outcome(s) assessed; longest follow-up
		Age: NR Gender: 61.6% male			Unclear Was ITT analysis undertaken: No	
Multiple interventions Wasserman et al. (2015) [101] Multi-site: 10 European countries	Inclusion: all students in participating classrooms Exclusion: None (although students who reported suicide attempts ever, or severe suicidal ideation in the past 2 weeks before the baseline assessment, and those with missing data regarding these two variables were excluded from the final analysis) Recruited from: Secondary schools (N = 168)	Whole sample N = 11,110 Mean age: 14.8 (SD:0.82) Range: 14–16) Gender: NR Treatment group 1 N = 2692 Mean age: 14.8 (SD: 0.82) Gender: 37% male Treatment group 2 N = 2721 Mean age: 14.8 (SD: 0.85) Gender: 40% male Treatment group 3 N = 2764 Mean age: 14.8 (SD: 0.8) Gender: 42% male Control group N = 2933 Mean age: 14.78 (SD: 0.89) Gender: 44% male	Psycho-educational (universal) component Youth Aware of Mental Health Programme (YAM), a universal intervention that aims to raise awareness of risk & protective factors associated with suicide, including knowledge of depression/anxiety and to enhance skills to manage stress, adverse life events & suicidal behaviors Length: 3 h role play session plus 2 × 1 h lectures Developed by: study authors Gatekeeper training (selective) component Question, Persuade, and Refer (QPR), a gatekeeper training module targeting teachers and other school personnel. Length: NR Developed by: Tompkins et al. (2010) ^j Screening (selective) component Screening by health professionals (ProfScreen) with referral of at-risk pupils. Length: NA Developed by: study authors Delivered by: Trained instructors	Active placebo: The control group was exposed to the same 6 educational posters as the YAM group. These included information about local health-care providers.	Random sequence generation method: Cluster stratified randomization using a random numbers table Allocation concealment method: NR Ascertainment of SH repetition: Self-report Outcome assessor blinding: NA Less than 15% drop-out rate: No (26.3%) Was ITT analysis undertaken: No	SI (dichotomous): single item from five item Paykel Hierarchical Suicidal Ladder SA (dichotomous): single item from five item Paykel Hierarchical Suicidal Ladder Longest follow-up:12 months (not specified if post-test or post-intervention)

Notes: ED = Emergency Department; ITT = intention-to-treat; IQR = Interquartile Range; MA = meta-analysis; MH = mental health; NA = not applicable; NR = not reported; TAU = treatment as usual; SA = suicide attempt; SD = standard deviation; SH = self-harm; SI = suicidal ideation; SRB = suicide-related behavior.

^a Ross CP. School and suicide: Education for life and death. In RFW Diekstra & K Hawton (Eds.), *Suicide in adolescence*. Dordrecht: Martinus Nijhoff, 1987.

^b Excluded secondary publication: Randell BP et al. Immediate post-intervention effects of two brief youth suicide prevention interventions. *Suicide Life Threat Behav* 2001; 31(1): 41–61.

^c D'Zurilla TJ, Nezu AM. Development and preliminary evaluation of the Social Problem-Solving Inventory. *Psychol Assess* 1990; 2: 156–163.

^d Note: This study recruited participants from both schools and the community.

^e Van Orden KA, et al. Thwarted belongingness and perceived burdensomeness: construct validity and psychometric properties of the Interpersonal Needs Questionnaire. *Psychol Assess* 2012; 24:197–215.

^f Classified as BCI in the meta-analysis

^g Linehan MM. *Skills training manual for treating borderline personality disorder*. New York: Guilford Press, 1993.

^h Robinson WL, Case MH. *Leader manual for the Down with Drama course*. Unpublished Manual. DePaul University; Chicago: Illinois; 1995.

ⁱ Mufson L, et al. Effectiveness research: Transporting interpersonal psychotherapy for depressed adolescents (IPT-A) from the lab to school-based health clinics. *J Consult Clin Psychol* 2004; 7: 251–261.

^j Tompkins TL, et al. Does a gatekeeper suicide prevention program work in a school setting? Evaluating training outcome and moderators of effectiveness. *Suicide Life Threat Behav* 2010; 40: 506–15.

Table 4

Study characteristics: Non-randomized controlled trials conducted in educational or workplace settings (N = 16).

Study; country	Study design; level of evidence	Target population	Participants	Intervention description	Comparison condition	Risk of bias	Suicide related outcome(s) assessed; Longest follow-up	Results	Interpretation
Universal interventions									
Bailey et al. (2017) [110] Australia	Study design: Pre-test/post-test case series Level of evidence: IV	Inclusion: year 11 and 12 students at participating schools Exclusion: None Recruited from: Secondary schools (N = 3)	Whole sample: N = 129 Mean age: 16.7 (range 16–18) Gender: 53.5% male	Educational safeTALK Length: One 3-hour session Developed by: LivingWorks Delivered by: Trained instructors	NA	Adequately powered: No Outcome assessor blinding: NA Less than 15% drop-out rate: No (25.9%) Use of statistical testing to measure change at from pre-test to post-test: Yes	SI: Single item asking participants to indicate whether or not they were experiencing current suicidal thoughts. Longest follow-up: 1 month post-intervention	SI: NR	In comparison with Time 1 (the reference category), individuals at Time 2 had 0.53 times the odds of experiencing suicidal thoughts (95% CI = 0.20–1.36), and at Time 3 had 0.30 times the odds (95% CI = 0.10–0.91).
King et al. (2011) [102] USA	Study design: Pre-test/post-test case series Level of evidence: IV	Inclusion: Students at participating schools Exclusion: None Recruited from: Secondary schools (N participating = NR, but the program was implemented in 24 schools).	Whole sample: N = 1030 Mean age: 14.1 (SD: 0.79; range 14–18) Gender: 43.9% male	Educational Surviving the Teens® Suicide Prevention and Depression Awareness Program. Length: Four 50-min sessions Developed by: Study authors Delivered by: MH professionals	NA	Adequately powered: No power calculations provided. However, likely to be adequately powered for SI but not SA. Outcome assessor blinding: NA Less than 15% drop-out rate: No (59.6%) Use of statistical testing to measure change at from pre-test to post-test: Yes	SI: Single item asking participants to indicate whether or not they were currently seriously considering attempting suicide Longest follow-up: 3 months post-intervention	SI: Post-test Yes = 2.9% (n = 26) Follow-up Yes = 1.5% (n = 6)	There was no significant change in the number of students seriously considering suicide at post-intervention from pre-intervention ($\chi^2 = 0.837$, $p =$ 0.360). At 3-month follow-up, students were significantly less likely than at pre-test to be currently considering suicide ($p = 0.035$).
LaFromboise & Howard-Pitney, (1994) [93] USA	Study design: Non-randomized experimental trial Level of evidence: III-2	Inclusion: Students attending a Zuni secondary school Exclusion: None Recruited from: Secondary school (N = 1)	Whole sample: N = 83 Mean age: 15.6 SD/Range: NR Gender: 41% male Treatment group: NR Control group: NR	Educational The Zuni Life Skills Development Curriculum. Units included: information about suicide; suicide intervention skills; communication skills; coping with oppression; anger & stress management and goal setting. Length: Six units delivered across 28 lessons Developed by: Study authors Delivered by: Teachers	NR	Intervention developer: Study authors Adequately powered: Unclear Outcome assessor blinding: NA Less than 15% drop-out rate: No (25.3%) Use of statistical testing to measure change at from pre-test to post-test: Yes	SI: Suicide Ideation subscale of the Suicide Probability Scale (SPS) Longest follow-up: Post-intervention only	SI: Intervention mean (SD): 13.4 (NR) Control mean (SD): 16.8 (NR)	No between-group statistical analysis completed.
LaFromboise & Howard-Pitney, (1995) [94] USA	Study design: Non-randomized experimental trial Level of evidence: III-2	Inclusion: Freshman and junior students taking language arts classes at a Zuni secondary school Exclusion: None Recruited from:	Whole sample: N = 128 Mean age: 15.9 (Range: 14–19) Gender: 36% male Treatment	Educational The Zuni Life Skills Development Curriculum. Units: building self-esteem; identifying emotions & stress; communication & problem-solving skills; recognizing & eliminating self-destructive behavior; suicide information; suicide intervention training; goal setting	No intervention	Adequately powered: No Outcome assessor blinding: NA Less than 15% drop-out rate: No (23.4%) Use of statistical testing to	SRB: SPS Longest follow-up: 8 months post-baseline	SRB: Intervention mean (SD): 54.3 (SD: 11.6) Control mean (SD): 58.9 (SD: 13.0)	The treatment group was less suicidal after taking part in the curriculum than the control group, $t [61] = 1.45$, p < 0.07.

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Table 4 (continued)

Study; country	Study design; level of evidence	Target population	Participants	Intervention description	Comparison condition	Risk of bias	Suicide related outcome(s) assessed; Longest follow-up	Results	Interpretation
Vieland et al. (1991) [103] USA	Study design: Non-randomized experimental trial Level of evidence: III-2	Secondary school (N = 1) Inclusion: Ninth grade students from participating schools Exclusion: Schools excluded if they had ever received a suicide prevention program Recruited from: Secondary schools (N = 4)	group: N = 69 Age/gender: NR Control group: N = 59 Age/gender: NR Whole sample: N = 381 Mean age: 15.8 Gender: NR Treatment group: N = 174 Mean age: 15.8 (SD: 0.64) Gender: 45% male Control group: N = 207 Mean age: 15.8 (SD: 0.59) Gender: 51% male	Length: Seven units delivered 3 times a week over approx. 30 weeks Developed by: Study authors Delivered by: Teachers Educational In-class presentation. Emphasized support networks in alleviating stress, confronting one's peers, and community resources. Length: 1.5 h Developed by: Unclear Delivered by: Teachers	No intervention	measure change at from pre-test to post-test: Yes Adequately powered: No Outcome assessor blinding: NA Less than 15% drop-out rate: Unclear Use of statistical testing to measure change at from pre-test to post-test: No	SA: Single item asking participants to indicate whether or not they had made a first suicide attempt Longest follow-up: 18 months post-baseline	SA: Intervention: Yes = 2.5% Control: Yes = 2.7%	There was no evidence that the program had an effect on suicide attempt rates.
Selective interventions Hazell & Lewin (1993) [104] Australia	Study design: Post-test case series Level of evidence: IV	Inclusion: Students who had been exposed to the suicide of a peer Exclusion: None Recruited from: Secondary schools (N = 2)	Whole sample: N = 126 Mean age/gender: NR Treatment group: N = 63 Age/gender: NR Control group: N = 63 Age/gender: NR	Therapeutic One session of group counseling provided at school within 7 days of a student suicide. Following the session, school staff were debriefed & arrangements made to follow-up high risk students. Length: 90 min Developed by: Study authors Delivered by: MH professionals	Unclear	Adequately powered: NR Outcome assessor blinding: NA Less than 15% drop-out rate: Unclear Use of statistical testing to measure change at from pre-test to post-test: Yes	SA: Hospitalization for SA assessed using Youth Self Report (YSR) version of the Child Behavior Checklist (CBCL) SH: Incidence of current suicidal behavior - YSR CBCL SI: % of group currently experiencing suicidal ideation - YSR CBCL Longest follow-up: Post-intervention only	SA: Intervention: 1.6% Control: 0.0% SH: Intervention: 21.0% Control: 19.0% SI: Intervention: 14.5% Control group: 19.0%	There were no differences between groups on SA, SH or SI as assessed by Pearson X2
McDaniel et al. (1990) [121] USA	Study design: Interrupted time series with a control group Level of evidence: III-2	Inclusion: US Navy instructors Exclusion: None Recruited from: Navy training command (N = 1)	NR	Training sessions for instructors in US Navy training command. Focused on how instructors can identify signs of distress and risk in their students, how to intervene and how to get help. Length: 3 × 1 h Developed by: Unclear Delivered by: MH professionals	"Operational command" - less than 10 miles away from training command and about the same size but no training.	Adequately powered: NR Outcome assessor blinding: NA Less than 15% drop-out rate: Unclear Use of statistical testing to measure change	SA: Average monthly rate of SA (obtained from official sources) Longest follow-up: NA	SA: Post-test Intervention rate: 9.4 Control rate: 1.8	There was a declining trend in the suicide attempt rate in the intervention group. At post-test, the average monthly suicide attempt rate was significantly higher in the intervention group, $p < 0.001$.

					at from pre-test to post-test: Yes				
Indicated interventions									
Biddle et al. (2014) [105] USA	Study design: Post-test case series Level of evidence: IV	Inclusion: Students demonstrating SRB Exclusion: NA Recruited from: Secondary schools (N = 619)	N = 18,445 Mean age: NR (Range: 13–21) Gender: NR	Therapeutic <i>Student Assistance Program (SAP)</i> : Identify individual student problems & recommend interventions. Participants are students referred to the SAP who accessed the recommended services. Length: NA Developed by: Commonwealth Student Assistance Program Interagency Committee, Pennsylvania. Delivered by: Trained school staff	No intervention	Adequately powered: Yes Outcome assessor blinding: NA Less than 15% drop-out rate: Unclear Use of statistical testing to measure change at from pre-test to post-test: Unclear	Suicide: Number of suicides and suicide rate per 100,000 students Longest follow-up: NA	Suicide: Post-test Intervention N (rate): 9 (65.2) Control N (rate): 6 (129.25)	The difference in suicide rates was not statistically significant.
Eggert et al. (1995) [106]; (1999) ^a [36], USA	Study design: Post-test case series Level of evidence: IV	Inclusion: Students reporting: SA; SI; moderate-serious depression; specific levels of alcohol or other drug use, polyuse, or drug use control problems Exclusion: None Recruited from: Secondary schools (N = 5)	Whole sample: N = 105 Mean age/gender: NR PGC I group: N = 36 Mean age: 16.19 (SD: 0.92) Gender: 41.7% male PGC II group: N = 34 Mean age: 15.82 (SD: 1.11) Gender: 37.1% male Control: N = 35 Mean age: 15.57 (SD: 1.01) Gender: 45.7% male	Psycho-education Personal growth classes (PGCs): Incorporated [1] group work; [2] weekly monitoring of activities targeting changes in mood management, school performance and attendance, and drug involvement; and [3] life skills training in self-esteem enhancement, decision making, personal control (skills training in anger, depression, and stress management), and interpersonal communication. Length: (PGC I): One semester - 5 months or 90 class days; (PGC-II): Two semesters - 10 months or 180 class days. Developed by: Study authors Delivered by: Trained school staff	Enhanced TAU: Assessed for suicide 'potential'.	Adequately powered: NR Outcome assessor blinding: NA Less than 15% drop-out rate: Unclear Use of statistical testing to measure change at from pre-test to post-test: Yes	SRB: <i>The Brief Suicide Risk Behavior Scale</i> ; A 5-item scale measuring the frequency of suicide thoughts, direct and indirect suicide threats, and suicide attempts Longest follow-up: 10 months post-intervention	SRB: Post-test PGC1 mean (SD): 0.27 (0.55) PGC2 mean (SD): 0.66 (0.85) Control mean (SD): 0.55 (0.65) Follow-up PGC1 mean (SD): 0.56 (0.93) PGC2 mean (SD): 0.66 (0.79) Control mean (SD): 0.23 (0.37)	There was no significant difference in suicide risk behaviors between the groups. There was a significant decline in suicide risk behaviors for all three groups (F Linear [1,102] = 104.14, $p < 0.001$) revealed a significant decline for all three groups.
Joffe (2008) [118] USA	Study design: Interrupted time series with a control group Level of evidence: III-2	Inclusion: Students with a SA or suicide 'threat' Exclusion: NA Recruited from: University (N = 1)	Student population Treatment location: 1980–1983: 139,384 1984–1990: 249,812 Control location: 1980–1983: 1,244,469 1984–1990: 1,807,968	Policy Implementation of a policy requiring any student who made a suicide threat or attempt to receive 4 individual sessions of professional assessment, the first which occurred within a week of the incident. Length: NR Developed by: Counseling Center, University of Illinois Delivered by: MH professionals	Data collected from 11 other universities	Adequately powered: NR Outcome assessor blinding: NA Less than 15% drop-out rate: Not Reported Use of statistical testing to measure change at from pre-test to post-test: Yes	Suicide: Deaths by suicide per 100,000 enrolled students per year Longest follow-up: NA	Suicide: Post-test Intervention rate: 2.0 Control rate: 8.68	The treatment group had a 74.7% reduction in the suicide rate, compared to an increasing suicide rate in the comparison group, z score = 5.90, $p < 0.05$
Lerner & Clum (1990) [117] USA	Study design: Non-randomized, experimental trial Level of evidence: III-2	Inclusion: Students with SI Exclusion: Psychosis, substance abuse	Whole sample N = 18 Mean age: 19.17 (SD:	Therapeutic group problem solving therapy Length: 10 sessions over 5–7 weeks Developed by: Based on D'Zurilla and Goldfried (1971) ^b	Active placebo: empathetic listening, sharing experiences with the group.	Adequately powered: No Outcome assessor blinding: NA Less than 15%	SI: Modified Scale for Suicidal Ideation (MSSI) Longest follow-up: 3 months post-intervention	SI: Post-test Intervention mean (SD): 5.8 (7.0) Control mean	There was no significant difference in suicidal ideation between the groups at both time points (T2: F value < 1; T3: F value = 1.87)

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Table 4 (continued)

Study; country	Study design; level of evidence	Target population	Participants	Intervention description	Comparison condition	Risk of bias	Suicide related outcome(s) assessed; Longest follow-up	Results	Interpretation
		Recruited from: University (N = 1)	1.38; Range: 18–24) Gender: 22% male Treatment group N = 9 Mean age: 18.78 (SD: 0.83; Range: NR) Gender: 11% male Control group N = 9 Mean age: 19.56 (SD: 1.74) Gender: 33% male	Delivered by: MH professional		drop-out rate: Yes (0.0%) Use of statistical testing to measure change at from pre-test to post-test: Yes		(SD): 5.3 (9.2) 3-month follow-up Intervention mean (SD): 4.7 (3.4) Control mean (SD): 10.6 (8.8)	
Robinson J et al. (2016) ^c [112], Australia	Study design: Pre-test/post-test case series Level of evidence: IV	Inclusion: Students who presented to the school counselor with SI in the past month Exclusion: ID, psychosis, inability to speak English Recruited from: Secondary schools (N = 11)	Whole sample: N = 32 Mean age: 15.6 (Range: 14–17) Gender: 12.5% male	Therapeutic Reframe-IT. Individual online suicide-specific CBT: eight 20-min modules incorporating standard CBT approaches commonly used with young people. Length: eight 20-min modules Developed by: Study authors Delivered by: self-directed	NA	Adequately powered: No Outcome assessor blinding: NA Less than 15% drop-out rate: No (38.2%) Use of statistical testing to measure change at from pre-test to post-test: Yes	SI: Suicidal Ideation Questionnaire (SIQ) Longest follow-up: Post-intervention only	SI: Pre-test Mean (SD): 3.2 (1.6) Post-test Mean (SD): 1.5 (1.3)	There was a statistically significant decrease in SI from pre to post-test, with a moderate effect size, $t = 6.2$; p < 0.0005
Multi-modal interventions									
Aseltine et al. (2007) ^d [107], USA	Study design: Pseudo-RCT Level of evidence: III-1	Inclusion: Students at participating schools Exclusion: None Recruited from: Secondary schools (N = 9)	Whole sample N = 4133 Mean age/range: NR Gender: 50.0% male Treatment group: 2039 Control group: 2094	Psycho-education & screening Signs of Suicide (SOS). <i>Universal educational component:</i> Video & discussion guide depicting signs of suicidality & depression and recommended ways to respond. <i>Selective component:</i> Screening to identify students at risk. Length: Video = 17 min Developed by: Screening for Mental Health Inc. Delivered by: NR	Attended class as usual & received the program after the study period	Adequately powered: No power calculations provided. However, likely to be adequately powered for SI but not SA. Outcome assessor blinding: NA Less than 15% drop-out rate: Unclear Use of statistical testing to measure change at from pre-test to post-test: No	SA: Item from the Youth Risk Behavior Survey (YRBS): During the past 3 months did you actually attempt suicide? SI: Item from the YRBS: During the past 3 months did you ever seriously consider attempting suicide? Longest follow-up: 3 months post-intervention	SA: Post-test % answered “Yes” Intervention: 3.0% Control: 4.6% SI: Post-test % answered “Yes” Intervention: 10.1% Control: 11.5%	Participants in the treatment group were 40% less likely to report a SA in the past 3 months compared with participants in the control group, $\beta = -0.47$ (SE 0.16), $p = 0.0075$ (OR = 47%) There was no significant effect of the SOS program on SI, $\beta = -0.53$ (SE = 1.01), $p > 0.05$
Shelif et al. (2016) [120] Israel	Study design: Interrupted time series with a	Inclusion: Active duty mandatory service military	Whole sample: N = 1,171,359	Multiple Israeli Defense Force Suicide Prevention Program. Includes: means restriction,	Cohort inducted into the IDF prior to the	Adequately powered: Yes Outcome assessor	Suicide: Total number of suicides & average number per year	Suicide: Pre-intervention (2006–2012): N	Trend analysis showed lower suicide rates in the cohort after intervention, Hazard ratio = 0.48 (95%CI:

	control group Level of evidence: III-2	personnel who served between 1992 and 2012 Exclusion: subsection of the population that did not represent the regular mandatory service soldiers Recruited from: Israeli Defense Force	Mean age: 19.0 (Range: 17–24) Gender: 53.4% male Treatment group: N = 405,252 Mean age: 19.0 Gender: 55.2% male Control group: N = 766,107 Mean age: 19.0 Gender: 15.4% male N = 3244	improved screening & management of suicidal soldiers, psycho-education and gatekeeper training. Length: NA Developed by: Based on Knox et al. (2003) ^e Delivered by: NR	implementation of the intervention (1992–2005)	blinding: NA Less than 15% drop-out rate: NR Use of statistical testing to measure change at from pre-test to post-test: Yes	Longest follow-up: NA	= 344; 24.6 per year Post-intervention (1992–2005): N = 89; 12.7 per year	0.37–0.60)
Silverstone et al. (2015) [108]; (2017) [37] Canada	Study design: Pre-test/post-test case series Level of evidence: IV	Inclusion: All secondary school students, plus targeted intervention for students with SI (with or without SA) Exclusion: None Recruited from: Secondary schools (N = 5)	Mean age: NR (Range: 10–19) Gender: 51.7% male	Therapeutic & screening Empowering a Multi-modal Pathway Toward Healthy Youth (EMPATHY): Universal CBT for all students in years 7 and 8; screening for all students; rapid intervention, guided online CBT for those identified as being at-risk. Length: Universal CBT = 8–16 sessions; online CBT = NR Developed by: Study authors Delivered by: MH professionals	NA	Adequately powered: Yes Outcome assessor blinding: NA Less than 15% drop-out rate: No (24.2%) Use of statistical testing to measure change at from pre-test to post-test: Yes	SI: N at “High risk” (“thought you were better off dead” more than half the days in past 2 weeks) or “medium risk” (“thought you were better off dead” several days in past 2 weeks) of suicide Longest follow-up: 15 months post-baseline	SI: 12-week follow-up High risk N = 30 Medium risk N = 19 Actively suicidal N = 49 15-month follow-up High risk N = 16 Medium risk N = 21 Actively suicidal N = 37	Less students were at “high” or “medium” risk of suicide at follow-up compared to baseline (significance testing not reported). At 15-month follow-up, significantly less people were “actively suicidal” (high or medium risk) than at baseline, $p < 0.001$
Zenere & Lazarus, (1997) [109] USA	Study design: Interrupted time series without a control group Level of evidence: III-3	Inclusion: All public secondary school students in Miami, Florida Exclusion: NA Recruited from: Secondary schools (N = 300)	NR	Multiple Suicide Prevention and School Crisis Management Program: Comprising psycho-education, postvention, school-based crisis teams, staff training, crisis hotline. Length: NA Developed by: Dade County Public Schools Department, Florida. Delivered by: MH professionals & teachers.	Cohort of students at beginning of implementation of the SPSCMP (1989–1990)	Adequately powered: No power calculations provided. However, likely to be adequately powered for SI but not SA. Outcome assessor blinding: NA Less than 15% drop-out rate: NR Use of statistical testing to measure change at from pre-test to post-test: No	SI: Number of suicidal ideations, obtained via hotline reports SA: number and rate of suicide attempts, obtained via hotline reports Longest follow-up: NA	SI: Pre-test (1989–1990) N = 641 Post-test (1993–1994) N = 640 SA: Pre-test N = 243; 87/100,000 Post-test N = 95; 31/100,000	The number of suicidal ideations among students fluctuated during the data collection period, initially showing a decrease in activity before returning to previous levels. The rate of suicide attempts decreased

Notes: ED = Emergency Department; ID = Intellectual Disability; ITT = intention-to-treat; IQR = Interquartile Range; MH = mental health; NA = not applicable; NR = not reported; TAU = treatment as usual; SA = suicide attempt; SD = standard deviation; SH = self-harm; SI = suicidal ideation; SRB = suicide-related behavior.

^a 1999 is a correction; excluded secondary publication: Thompson EA, et al. Mediating effects of an indicated prevention program for reducing youth depression and suicide risk behaviors. *Suicide Life Threat Behav* 2000; 30(3): 252–71.

^b D’Zurilla T, Goldfried M. Problem solving and behavior modification. *J Abnorm Psychol* 1971; 78: 107–126.

^c Excluded secondary publications: Robinson J, et al. The safety and acceptability of delivering an online intervention to secondary students at risk of suicide: findings from a pilot study. *Early Interv Psychiatry* 2015; 9(6): 498–506.; Hetrick S, et al. Does cognitive behavioural therapy have a role in improving problem solving and coping in adolescents with suicidal ideation? *Cognitive Behaviour Therapist* 2014; 7.

^d Excluded secondary publication: Aseltine RH, DeMartino R, An Outcome Evaluation of the SOS Suicide Prevention Program. *American Journal of Public Health* 2004; 94(3): 446–451.

^e Knox KL, et al. Risk of suicide and related adverse outcomes after exposure to a suicide prevention programme in the US Air Force: cohort study. *BMJ* 2003; 327(7428):1376.

Table 5
Interrupted time series and ecological studies in community settings (N = 15).

Study; country	Study design; level of evidence	Target region/population; comparison	Intervention description	Time period	Risk of bias	Outcome/data source	Rates per 100,000	Interpretation
Universal: means restriction								
Beautrais et al. (2006) [125] New Zealand	Study design: Interrupted time series without a control group Level of evidence: III-3	Target region/population: NA Comparison: None	Firearms legislation introduced in 1992 mandating license to own a firearm.	1985–1992: pre-legislation; 1993–1996: implementation; 1997–2002: post-implementation.	Were data collected at multiple time points? Yes Was the intervention likely to affect data collection? No: Data collection from official mortality data at a national level	Suicide: Mean annual age-specific suicide rates by all methods and by firearm for persons aged 15–24 years, obtained through New Zealand Health Information Service (NZHIS)	Suicide: All: Unclear Firearm: Unclear	There was a 66% decrease in the mean annual rate of firearm-related suicide ($B = -1.09$; $SE = 0.24$; $p < 0.001$). There was a decrease in the fraction of all suicides accounted for by firearm-related suicides ($p < 0.0001$). There was no significant decrease in overall rates of suicide ($b = 0.08$; $SE = 0.10$; $p = 0.39$).
Caron (2004) [126] Canada	Study design: Interrupted time series without a control group Level of evidence: III-3	Target region/population: Northern Quebec Comparison: None	Firearms legislation introduced in 1992 mandating firearm owners to safely store their firearms.	1986–1991: pre-legislation; 1992–1996: post-legislation	Were data collected at multiple time points? Yes Was the intervention likely to affect data collection? No: Data collection from official mortality dataset	Suicide: Age-specific suicide rates by all methods and by firearm for under 25 age group obtained through the Quebec Coroner's office.	Suicide: All: NR Firearm: NR	There was a 38% decrease in firearm suicides (significance = NR). There was a 69% increase in the overall suicide rate ($X^2 = 22.09$, $df = 1$, $p < 0.001$).
Cheung and Dewa (2005) [128] Canada	Study design: Interrupted time series without a control group Level of evidence: III-3	Target region/population: NA Comparison: none	Restrictive firearms regulations - Bill C-17 enacted in 1991.	1979: pre-implementation 1999: post-implementation	Were data collected at multiple time points? Yes Was the intervention likely to affect data collection? No: data collection from official mortality dataset.	Suicide: Age-specific suicide rates for youth between 15 and 19 years by firearm, overdose, hanging, and total, and percentage of suicides by each particular method, obtained from data collected by the Coroner's office.	Suicide: All: Unclear Firearm: 1979: Unclear 1999: 3.0 Overdose: 1979: 1.2 1999: 0.2 Hanging: 1979: 2.6 1999: 7.8	The findings suggest a substantial decrease in firearm-related suicides but no decrease in the overall suicide rate (significance = NR).
Leenaars & Lester (1997) [131] Canada	Study design: Interrupted time series without a control group Level of evidence: III-3	Target region/population: NA Comparison: None	Gun control legislation introduced in 1977 (Bill C-51).	1969–1976: pre-legislation 1978–1985: post-legislation	Were data collected at multiple time points? Yes Was the intervention likely to affect data collection? No: data collection from official sources.	Suicide: Suicides rates by firearm and by all methods, and percentage of total suicide rate by firearm, in persons aged 15–24, obtained from Statistics Canada and supplemented by personal communications.	Suicide: All: Pre-legislation: 12.57 Post-legislation: 16.11 Firearm: Pre-legislation: 5.89 Post-legislation: 7.12	There was a significant increase in the mean firearm suicide rate and mean total suicide rate from pre-legislation to post-legislation ($p < 0.05$). There was no statistically significant change in the percentage of all suicides that were by firearm.
Lubin et al. (2010) [132] Israel	Study design: Interrupted time series without a control group Level of evidence: III-3	Target region/population: Defense Force personnel Comparison: None	Rule prohibiting soldiers from taking home service weapons on the weekend.	T1: 2003–2005 T2: 2007–2008	Were data collected at multiple time points? No Was the intervention likely to affect data collection? No: Israeli Defense Force suicide data.	Suicide: Average number of suicide deaths per year; firearm suicides on weekends; firearm suicides on weekdays in soldiers aged 18–21: data source not specified.	Suicide: All: NR Firearm suicides on weekends: NR Firearm suicides on weekdays: NR	Following policy change, suicide rates decreased significantly by 40% ($t = 3.35$, $p = 0.04$). Most of this decrease was due to decrease in suicide using firearms over the weekend ($t = 17.44$, $p < 0.001$). There was no significant change in rates of suicide on weekdays
Niederkrötenhaler et al. (2009) [135] Austria	Study design: Interrupted time series without a control group Level of evidence: III-3	Target region/population: NA Comparison: None	1997 revision of firearm laws to harmonize with EU regulations	T period 1–1986–1987 T period 2 - 1987–2006	Were data collected at multiple time points? Yes Was the intervention likely to affect data collection? No: National mortality data.	Suicide: Suicide rates by firearm and all methods in 10–19 year-olds, obtained from Statistics Austria.	Suicide: All: NR Firearm: Unclear	There was a temporary increase in firearm suicides, followed by a continuous decrease (adjusted -0.20 95% CI -0.33 to -0.07 ; $p = 0.003$). On the whole, firearm suicide rates after the firearm legislation reform were significantly lower than before the reform. The adjusted model showed no changes in total suicide rates associated with the reform (adjusted 0.017 95%CI 0.04 – 0.074 ; $p = 0.533$).
Wheeler et al. (2009) [137] Multi-national (23 Countries of 35 with available suicide data from the WHO's Mortality Stratum A). Wheeler et al.	Study design: Interrupted time series without a control group Level of evidence: III-3 Study design: Target	Target region/population: NA Comparison: None	"Regulatory action" to restrict use of SSRIs Restriction of SSRIs	Pre-intervention: 1990–2003 Post-intervention: 2004–2006 Suicide deaths	Were data collected at multiple time points? Yes Was the intervention likely to affect data collection? No: National mortality data & Hospital admissions data. Were data collected at	Suicide: Suicide rates in 10–14 and 15–19 year-olds, obtained from WHO mortality database. Suicide: Mortality rates due to	Suicide: NR Suicide: Unclear	There was no evidence for an overall effect on the incidence of suicide of regulatory action regarding SSRIs for 15–19 year-olds ($p = 0.95$) or 10–14 year-olds ($p = 0.97$). There was no statistical evidence of

(2008) ^b [136], UK	Interrupted time series without a control group Level of evidence: III-3	region/population: Young people treated with SSRIs Comparison: None		Period 1–1993–2003 Period 2–2003–2005 Self-harm Period 1 1999–2003 Period 2–2003–2005	multiple time points? Yes Was the intervention likely to affect data collection? No: National mortality data & Hospital admissions data.	intentional self-harm or undetermined intent in 12–17 year-olds, obtained from Office for National Statistics. SA/SH: Hospital admissions per year due to intentional self-harm or undetermined intent in 12–17 year olds, obtained from the Department of Health Hospital Episode Statistics	Admissions for SA/SH: Males 1999: ~120 2005: ~120 Females 1999: 367 2005: 525	changes in trends in suicide rates between 1993 and 2005. The rate of hospital admissions remained relatively stable in males and steadily increased in females.
Multi-modal Ahmadi & Ytterstad (2007) [124] Iran	Study design: Interrupted time series with a control group Level of evidence: III-2	Target region/population: Young women and low SES in 2 cities Comparison: Sarpolzahab hospital (reference group)	Multi-modal: mix of passive and active strategies (not described). Key feature was psycho-education via videos	Pre-intervention - 1999–2000 Intervention - 2000–2003	Were data collected at multiple time points? No Was the intervention likely to affect data collection? Yes: Possible that those involved in data collection were not blinded to the intervention (suicide attempts).	SH: N (%) of total self-inflicted burn cases who was admitted in Gilangharb and Sarpolzahab hospitals during the baseline year to the study, during the study period, and the last year of the study period in persons aged 0–20 years.	SH: NR	No statistical analyses were performed on rates of self-immolation in youth.
Center for Disease Control (1998) [127] USA	Study design: Interrupted time series without a control group Level of evidence: III-3	Target region/population: Western Athabaskan tribe in rural New Mexico, USA Comparison: None	Multi-modal: gatekeeper training, outreach to families, immediate response and follow up for reported at-risk youth, community psychoeducation, and screening in services.	1988–1989: pre-implementation; 1990–1999: post-implementation	Were data collected at multiple time points? Yes Was the intervention likely to affect data collection? It is possible that those involved in data collection were not blinded to the intervention (suicide attempts). This was a not a problem with suicide deaths as this was obtained from official sources.	SRB: Rates of suicide acts for persons aged 15–19 (included completions and attempts) obtained via a surveillance form.	SRB: 1988–1989: 59.8 1990–1991: 8.9 1992–1993: 9.2 1994–1995: 17.6 1996–1999: 10.9	Although rates varied after implementation of the program, they remained substantially lower than before the program was initiated.
Cwik et al. (2016) [129] USA	Study design: Interrupted time series without a control group Level of evidence: III-3	Target region/population: Apache Indians Comparison: None	Multi-modal: implemented in 2006, included psychoeducation for students, gatekeeper training, and indicated interventions for suicidal young people.	2001–2006: pre-implementation 2007–2012: post-implementation	Were data collected at multiple time points? Yes Was the intervention likely to affect data collection? Yes: it is possible that those involved in data collection were not blinded to the intervention (suicide attempts).	Suicide: Suicide rates for persons aged 10–24 years, obtained via The Celebrating Life surveillance system (established by tribal resolution in 2001).	Suicide: Pre-test: 10–14 years: 17.1 15–19 years: 23.6 20–24 years: 151.9 Post-test: 10–14 years: 23.6 15–19 years: 101.9 20–24 years: 96.0	The suicide rate <i>increased</i> by 38% in 10–14 year-olds, and <i>decreased</i> by 5.5% in 15–19 year-olds and 36.8% in 20–24 year-olds.
Hacker et al. (2008) [130] USA	Study design: Interrupted time series without a control group Level of evidence: III-3	Target region/population: Somerville, MA, USA Comparison: Massachusetts	Multi-modal: implemented between 2003 and 2005, included local trauma response network, community wide vigil, school based counseling, hospital beds made available, outreach to suicide survivors to offer services, youth leadership programs, media reporting guidelines, community-wide education.	1994–2003: pre-intervention period. 2003–2005: Intervention period. 2005 onwards: post-intervention.	Were data collected at multiple time points? Yes Was the intervention likely to affect data collection? No: data collection from official sources.	Suicide: Data on suicide rates for 10–24 year-olds obtained via death certificate data (examined from 2001 to 2007, and then from 1994 for comparison), mortality data from Massachusetts Department of Public Health for (1994–2005). SA/SH: Data on suicide attempts for 10–24 year-olds obtained via self-inflicted injury data from Massachusetts Department of Public Health for (1994–2005), hospital discharge data (1996–2006), 911 dispatch call data (2004 onwards), teen health survey conducted in Somerville High School.	Suicide: 1994–1999: Somerville: 6.04 2000–2005: Somerville: 9.77 Massachusetts: 4.27 SA/SH: 2005 Somerville: 47.3 Massachusetts: 73.7 2006 Somerville: 53.2 Massachusetts: 74.8	Overall the data indicates a decrease in the rate of suicide and suicide attempts (significance = NR).

(continued on next page)

Table 5 (continued)

Study; country	Study design; level of evidence	Target region/population; comparison	Intervention description	Time period	Risk of bias	Outcome/data source	Rates per 100,000	Interpretation
May et al. (2005) [134] USA	Study design: Interrupted time series without a control group Level of evidence: III-3	Target region/population: Western Athabaskan Tribal Nation, New Mexico, USA Comparison: None	Multi-modal: Surveillance, screening/clinical interventions with extensive outreach in multiple settings, school-based prevention programs, community education for adults and youths, training of 'natural helpers.	Baseline - 1988–1989; then two yearly numbers and yearly averages until 2002	Were data collected at multiple time points? Yes Was the intervention likely to affect data collection? Unclear: program providers also main data collectors for non-fatal behaviors, participants may avoid reporting.	SI: Teen health survey SRB: Number of self-harm incidents (attempts and gestures combined) in 11 to 24 year-olds obtained from staff case conference notes and Indian Health Service records.	SRB: NR	There was a significant decline in the number of combined gestures and attempts in 19–24 year-olds (coeff = -765 , $p = 0.001$) and 11 to 18 year-olds (coeff = -0.517 , $p = 0.048$).
Multiple interventions Garraza et al. (2015) [139]; Walrath et al. (2015) [38] USA	Study design: Ecological Level of evidence: III-2	Target region/population: 466 counties, USA Comparison: 1161 counties not exposed to suicide prevention efforts	Multiple: Activities funded by the Garrett Lee Smith (GLS) Memorial Suicide Prevention Program, implemented between 2006 and 2009. Includes gatekeeper training, psychoeducation programs, screening, improved community partnerships and linkages to service, postvention programs, and crisis hotlines.	At least 1 NSDUH respondent between 2008 and 2011, suicide mortality between 2007 and 2010	Were data collected at multiple time points? No Was the intervention likely to affect data collection? No: Data collection from official sources.	SA: Suicide attempt rates for each county following the implementation of the GLS program for the population that was approximately 16 to 23 years of age during implementation, obtained via self-report from the NSDUH between 2008 and 2011 Suicide: Suicide rates for persons aged 10–24 years between 2007 and 2010, obtained from the National Vital Statistics System.	SA: NR Suicide: NR	Suicide attempts: Counties implementing GLS program activities had significantly lower suicide attempt rates among youths 16 to 23 years of age in the year following implementation of the GLS program than did similar counties that did not implement GLS program activities (4.9 fewer attempts per 1000 youths [95%CI, 1.8–8.0 fewer attempts per 1000 youths]; $p = 0.003$). There was no evidence of longer-term differences in suicide attempt rates. Suicide deaths: Counties implementing GLS training had significantly lower suicide rates among the population aged 10–24 years in the year after GLS training than similar counties that did not implement GLS training (1.33 fewer deaths per 100,000; $p = 0.02$). No evidence of an effect beyond one year after training implementation. Suicide rates: Suicide prevention programs have a negative impact on the suicide rate (Males: -1.33 , SE 0.5; $p < 0.05$; Females -0.276 SE.08, $p < 0.05$)
Matsubayashi & Ueda (2011) [133] Multi-national	Study design: interrupted time series with a control group Level of evidence: III-2	Target region/population: 21 OECD nations Comparison: 10 OECD countries without a national suicide prevention program	Multiple: National prevention programs - specific interventions not specified or analyzed.	1980–2004 One time period, statistical models include date of implementation of suicide prevention program - varies for each country	Were data collected at multiple time points? No Was the intervention likely to affect data collection? No: National mortality data.	Suicide: Suicide rates in under 25 year-olds, obtained via the WHO mortality database	Suicide: 20.901 (mean total rate)	

Notes: NSDUH = National Survey on Drug Use and Health; NA = not applicable; NR = not reported; OECD = Organization for Economic Co-operation and Development; WHO = World Health Organization; SA = suicide attempt; SE = standard error; SES = socio-economic status; SH = self-harm; SI = suicidal ideation; SRB = suicide-related behavior.

^a Defined as at least twice before or at least twice after implementation of the intervention.

^b Note: it is likely that this study is a subset of the data included in Wheeler et al (2009).

Table 6
Non-randomized experimental trials in community settings (N = 1).

Study; country	Study design; level of evidence	Target population	Participants	Intervention description	Comparison condition	Risk of bias	Suicide related outcome(s) assessed; Longest follow-up	Results	Interpretation
Allen et al. (2017) USA [138]	Study design: Non-randomized experimental trial Level of evidence: III-2	Inclusion: Young people living in Yup'ik communities Exclusion: None Recruited from: Yup'ik communities in southwest Alaska	Whole sample N = 124 Mean age: NR Gender: NR Treatment group N = 128 Mean age (SD): 14.24(1.72) Gender: 43% male Control group N = 74 Mean age (SD): 14.62(1.82) Gender: 73% male	Qungasvik: A cultural intervention based in a local, Indigenous theory of personal and community change. Modules are individual, family, or community level. Length: one or more 1–3 h sessions. Developed by: study authors Delivered by: different cultural experts.	Community 2 – exposed to less-intensive intervention.	Intervention developer: Rasmus et al. (2014) ^a Adequately powered: NR Was SH a pre-determined outcome: No Outcome assessor blinding: NR Less than 15% drop-out rate for DSH at post-intervention: Unclear Use of statistical testing to measure change from pre-test to post-test: Yes	SI: "A Way to Live a Very Good, Beautiful Life"—Reasons for Life (RL): a cultural adaptation and strengths-based extension of the Brief Reasons for Living Inventory for Adolescents. Longest follow-up: post-intervention	SI: Post-intervention: Intervention mean (SD): 22.42 (4.99) Control mean (SD): 62.98 (17.49)	The treatment group showed a greater reduction in SI (indicated by increased scores on the RL) than the control group (d = 0.28, p < 0.05).

Notes: NR = not reported; SD = standard deviation; SI = suicidal ideation.

^a Rasmus SM, et al. Creating Qungasvik (a Yup'ik intervention "toolbox"): Case examples from a community-developed and culturally-driven intervention. Am J Community Psychol 2014; 54: 140–152.

One RCT in this category was not included in the meta-analysis. This investigated the impact of Parent-Adolescent CBT [72]; authors reported reduced suicidal ideation in both groups during active and maintenance treatment and at follow-up.

3.3.1.3. Study Quality. The majority of these studies used random sequence generation [40–51,53–56,58,59,61,62,64–71] (k = 28; 84.8%) and 21 (60.6%) used adequate allocation concealment strategies [42–46,49–51,53–55,58,59,61,62,64–67,69,70]. Of the 25 studies that assessed outcomes via interview, 13 (52.0%) reported assessor blinding [43–45,49,51,53,55,56,59,61,64,65,70]. Thirteen studies reported conducting intention-to-treat (ITT) analysis [42,46,48,53,54,56,58,59,61,64,65,69,72]. One study did not use ITT, but conducted a sensitivity analysis to assess the robustness of the findings [70]. Nineteen (57.6%) reported less than 15% drop out and were classed as low risk for the purpose of meta-analysis [41,43–46,49–51,53,54,56,58,61,63–69].

3.3.2. Other Study Designs

3.3.2.1. Study Description. All nineteen studies in this category tested indicated therapeutic interventions. The majority employed a pre-test/post-case series study design (k = 11; 57.9%) [73–83]. Sixteen (84.2%) recruited participants from community mental health services or hospitals, including inpatient and emergency department settings [73–80,83–91]. Interventions included DBT, CBT, and brief contact interventions. Sixteen (84.2%) of the studies in this category had a mean age of 18 or younger. Please see Table 2.

3.3.2.2. Study Efficacy. Two of the five studies testing a CBT-based intervention reported reductions in suicide-related behaviour [73,86], and three reported reductions in suicidal ideation [73,75,76]. Five of the six studies testing DBT reported reductions in suicide-related behaviour [74,77,79,80,88], and four reported reductions in suicidal ideation [74,77,84,88]. Two of the three studies testing family-based interventions reported reductions in suicidal ideation [83,89], and one reported a reduction in suicide attempts [89]. One study reported a reduction in the proportion of young people reporting a suicide attempt following exposure to a crisis intervention program [78], and one reported reduced suicidal ideation following telephone counseling [81]. One study tested a brief contact intervention and reported no between-group differences [85]. A study of a problem solving intervention reported a reduction in the proportion of participants reporting suicide attempts in the treatment group compared to controls [87]. Finally, a study testing an intervention for American Indians reported reductions in suicidal ideation over time [82]. Significance testing was not always conducted or reported for studies in this category.

3.3.2.3. Study Quality. Only seven studies had dropout rates of less than 15% [73,78,81,83,84,86,89]. All but one [89] were either underpowered or the adequacy of the sample size could not be determined. Eight studies used a comparison group [84–91]. Three assessed outcomes using interview-rated measures [87,90,91], and only one reported that outcome assessors were blinded to treatment allocation [91]. Fifteen studies (78.9%) conducted statistical testing to measure change from baseline [73–75,77,79–86,88,89,91].

3.4. Studies Conducted in Educational and Workplace Settings

Thirty-one studies recruited participants from educational or workplace settings; of these 21 (67.7%) were conducted in schools [92–112], seven (22.6%) in universities [113–119], two (6.5%) in military-based workplace settings [120,121], and one (3.2%) from both schools and public places in the community [122]. Twenty-one (67.7%) had a mean participant age of 18 years or younger, eight studies (25.8%) had

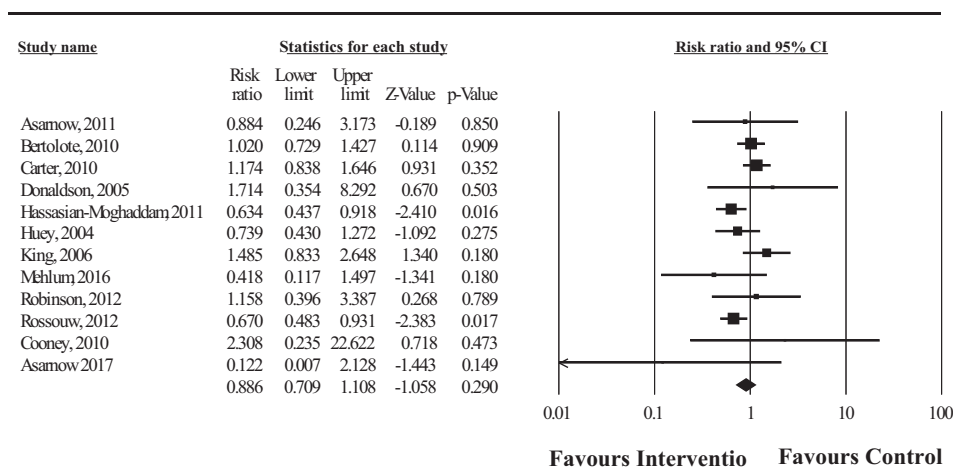


Fig. 2. Random effects risk ratio and 95% confidence interval (CI) for clinical interventions at the post-intervention assessment.

a mean age over 18, and in two studies (6.5%) the mean age could not be determined. Fifteen (48.4%) were RCTs.

3.4.1. Randomized Controlled Trials

3.4.1.1. Study Description. Three of the RCTs tested universal interventions [92,113,119], nine tested indicated interventions [95–98,114–116, 122,123], and three tested multi-modal or multiple interventions [99–101]. Studies were either educational or therapeutic in nature, and four tested an internet-based intervention [113,119,122,123]. One large cluster-RCT tested three distinct interventions (workshops for students; gatekeeper training; and screening) [101]. Two multimodal studies combined a universal educational component with screening. Examples of control conditions in these studies included TAU e.g. an interview with a school counselor, enhanced TAU, e.g. weekly therapy, and placebo e.g. a video about unrelated health issues. Ten studies (66.7%) in this category included participants with a mean age of 18 or under. See Table 3.

3.4.1.2. Study Efficacy. Eleven RCTs reported data amenable to meta-analysis [92,98,99,101,113–116,119,122,123]. Together there were 13 individual intervention arms because one study tested three interventions (one brief contact intervention and two universal educational interventions) [101]. Two intervention arms were brief contact interventions, five were universal educational interventions, and six were psychological interventions. As above findings are presented according to the outcome assessed, with the primary outcome (self-harm) reported first, followed by suicidal ideation. No studies reported suicide as an outcome.

3.4.1.2.1. Self-harm Measured Dichotomously. Compared to control, there was evidence of an intervention effect on self-harm at post-intervention ($k = 3$, $RR = 0.31$, 95% CI 0.15 to 0.61, $I^2 = 0\%$) (Fig. 4) and at follow-up ($k = 3$, $RR = 0.63$, 95% CI 0.42 to 0.96, $I^2 = 0\%$) (Fig. 5).

3.4.1.2.2. Sensitivity and Subgroup Analysis. As there were only three studies in this category these analyses were not possible.

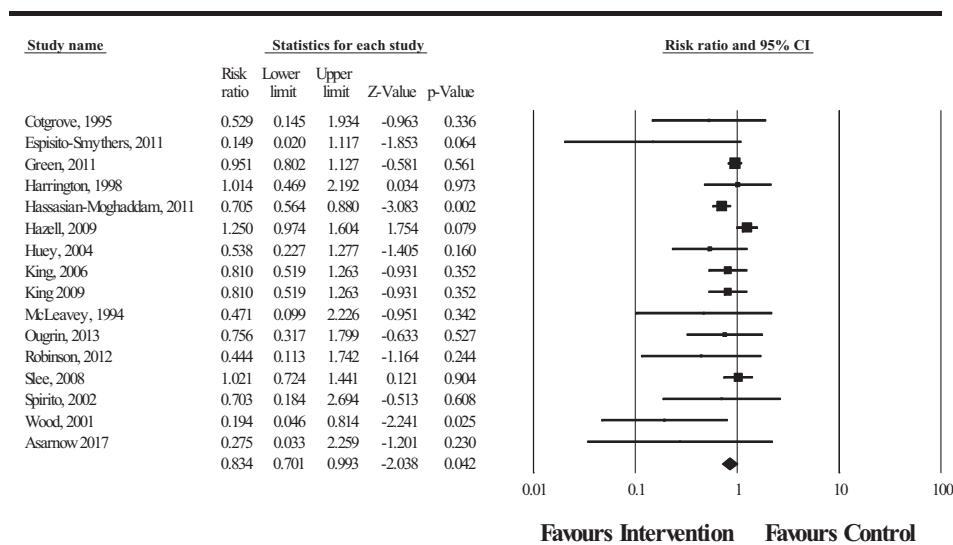


Fig. 3. Random effects risk ratio and 95% confidence interval (CI) for clinical interventions at the longest follow-up assessment.

3.4.1.2.3. Self-harm Measured Continuously. Compared to control, there was one study that reported continuous data post-intervention [115] with little evidence of an effect ($k = 1$, $SMD = -0.16$, 95% CI -0.61 to 0.30). No studies reported follow-up data for this outcome.

3.4.1.2.4. Suicidal Ideation Measured Dichotomously. Compared to control, there was little evidence of an effect at post-intervention ($k = 1$, $RR = 0.76$, 95% CI 0.50 to 1.16) or follow-up ($k = 2$ (4 intervention arms), $RR = 0.72$, 95% CI 0.51 to 1.03 , $I^2 = 0\%$).

3.4.1.2.5. Suicidal Ideation Measured Continuously. Compared to control, there was strong evidence of an effect of the intervention on suicidal ideation at post-intervention ($k = 7$, $SMD = -0.41$, 95% CI -0.57 to -0.24 , $I^2 = 15.2\%$). By follow-up, the effect was no longer significant ($k = 5$, $SMD = -0.21$, 95% CI -0.52 to 0.1 , $I^2 = 46.9\%$).

Four RCTs were not included in the meta-analysis. One tested a supportive intervention and found decreases in 'suicide risk behaviors' in treatment and control groups, but no between-group differences [95]. One examined a parent-specific intervention and found reductions over time in both groups, with greater reductions in the treatment group [96]. A group 'coping with stress course' tested with African-American adolescents was associated with a relative risk reduction in suicide in the intervention group compared to controls [97]. Finally, a multimodal intervention combining psycho-education and screening was associated with reduced suicidal ideation and behavior in intervention participants compared to controls [100].

3.4.1.3. Study Quality. Seven studies (46.7%) reported using random sequence generation techniques [95,99–101,116,122,123] and only three (20.0%) reported adequate concealment of treatment allocation [113, 122,123]. None of the studies in this category assessed primary outcomes using interviews, so outcome assessor blinding is not applicable. Six (40.0%) studies used ITT analysis [98,113,116,119,122,123]. One third ($k = 5$) had dropout rates of less than 15% [92,97,98,113,122].

3.4.2. Other Study Designs

3.4.2.1. Study Description. Of these 16 studies, four were non-randomized experimental trials [93,94,103,107,117], four were pre-test/post-test case series studies [102,108,110,112], three were post-test case series studies [104–106], and four employed an interrupted time series design [109,118,120,121]. The majority were conducted in school settings ($k = 12$; 75.0%), with two each (12.5%) conducted in university [117,118] and military settings [120,121]. Five studies tested universal educational programs [93,94,102,103,110], two evaluated selective interventions [104,121], five evaluated indicated interventions [105,106,112,117,118] and four evaluated multimodal interventions [93,94,102,103,107–110,120]. Two studies evaluated online interventions [108,112]. Eleven studies (68.8%) in this category had a mean participant age of 18 or under. See Table 4.

3.4.2.2. Study Efficacy. Of the five studies testing universal interventions, one reported a reduction in suicide-related behavior post-intervention [94], one reported a reduction in suicidal ideation post-intervention and at follow-up [110], and one reported a reduction at follow-up only [102]. Two studies tested selective interventions: one showed no effect of a counseling session delivered to school students bereaved by suicide [104] and the second reported a reduction in suicide attempts associated with a training intervention delivered to U.S. naval instructors [121].

Two of the five studies testing indicated interventions assessed suicide rates as the outcome of interest. The first found no impact of a therapeutic program among secondary school students [105]. The second examined the impact of a university suicide prevention policy and reported a reduction among the intervention group compared to increases among controls [118]. Of the remaining three studies of indicated interventions, only one therapeutic-based intervention was associated with a reduction in suicidal ideation from pre- to post-test [112].

Four studies tested a multimodal intervention. One was conducted in a workplace setting and reported lower suicide rates at post-intervention [120]. Two studies reported decreases in suicide attempts [107,109]. The final study examined the impact of a combined therapeutic and screening intervention and reported reductions in suicidal ideation at post-intervention and follow-up [108].

3.4.2.3. Study Quality. Only one study [117] reported an attrition rate of less than 15%. Three studies were adequately powered [105,108,120], and in another three, although no power calculations were provided, the sample size was sufficient to examine changes in suicidal ideation but not self-harm [102,107,109]. The majority of studies ($k = 12$; 75.0%) used statistical testing to measure change from pre- to post-test [93,94,102,104,106,108,110,112,117,118,120,121].

3.5. Studies Conducted in Community Settings

3.5.1. Study Description

Fourteen studies in this category (87.5%) were interrupted time series studies [124–137]; two (14.3%) utilized a control group [124,133]. One study was a non-randomized experimental trial [138] and one was an ecological study [139]. None of the community-based studies were RCTs. Eight (50.0%) evaluated means restriction approaches, five (31.3%) tested multimodal interventions [124,127,129,130,134] and two (12.5%) evaluated multiple interventions [133,139]. One non-randomized experimental trial [138] examined the impact of a cultural intervention among indigenous young people in Alaska.

3.5.2. Study Efficacy

Five of the six studies examining the impact of policies designed to restrict access to firearms reported decreases in the firearm suicide rate among young people [125,126,128,132,135], and one reported an

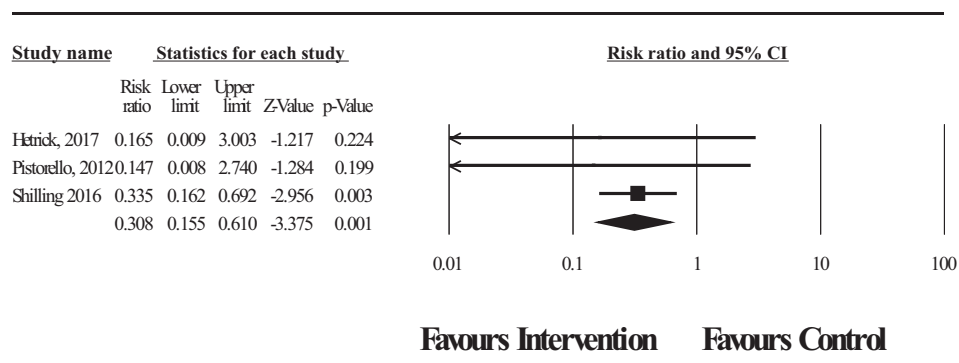


Fig. 4. Random effects risk ratio and 95% confidence interval (CI) for educational interventions at the post-intervention assessment.

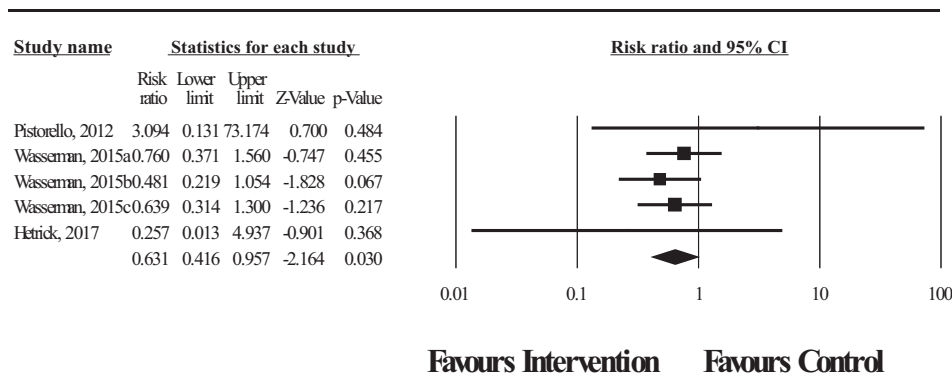


Fig. 5. Random effects risk ratio and 95% confidence interval (CI) for educational interventions at the longest follow-up assessment.

increase [131]. Only one reported a decrease in the overall youth suicide rate [132].

Two studies examined the impact of regulatory action to restrict use of antidepressants and found no evidence of an effect on suicide rates [136,137]. One of these studies also examined the impact of such regulatory action on rates of hospital admissions for self-harm and reported decreases in females only [136].

Three of the five studies evaluating multimodal interventions reported generally positive impacts on rates of suicide and/or suicide-related behaviour [127,130,134]. One study found the suicide rate decreased by 5.5% in 15–19 year-olds but increased by 38% in 10–14 year-olds [129]. Finally, one study evaluated the impact of an intervention targeting self-immolation in women; the authors reported a reduction in the number and percentage of self-immolation cases but did not report statistical significance [124].

One study evaluated multiple interventions delivered across different counties in the U.S. The interventions were associated with lower rates of suicide attempt [139] and suicide [38] but there was no evidence of a longer-term effect. Finally, a study evaluating the impact of government-initiated national suicide prevention programs across multiple nations reported decreases in suicide rates [133].

3.5.3. Study Quality

In 11 studies (73.3%), data were collected at multiple time points [125–131,134–137] and in 11 studies the intervention was deemed unlikely to impact data collection for the primary outcome of interest [125, 126,128,130–133,135–137,139].

4. Discussion

This review examined 99 individual studies of interventions designed to reduce suicide-related behaviors among young people. Samples were diverse, although few studies were conducted in low-to-middle income countries. Studies were conducted across a range of settings and tested a variety of intervention approaches, reflecting the spread of suicide prevention activity as recommended by current policy [7,1]. Less than half the studies were RCTs, which is unsurprising as the lack of RCTs in suicide prevention has been highlighted previously [24, 140]. Although not all intervention approaches, or intervention types, lend themselves to being tested this way, there remains a clear need for high-quality intervention studies in this field. In the majority of studies the mean age of participants was 18 or under (68.7%). In the clinical studies this was more prominent than in those conducted in educational settings (76.9% compared to 67.7%), suggesting that the findings from the clinical trials may be most applicable to young people aged 18 and under.

The number of intervention studies in youth suicide prevention has doubled in recent years, which is encouraging. However, many studies tested interventions originally designed for adults with little, or no,

adaption for young people [24]. This may partially account for the high rates of attrition in many of the studies reviewed. Adolescence and young adulthood are developmental periods requiring specific attention [141,142]. As such interventions that account for developmental stage and are both acceptable to, and ideally co-designed with, young people are necessary.

The meta-analysis showed little evidence that interventions reduced repetition of self-harm at post-intervention in clinical settings. Whilst there was some evidence for reduced repetition of self-harm at follow-up, this effect disappeared after removing low-quality studies; as such these findings should be interpreted with caution. There may be a small effect on frequency of self-harm measured continuously. It is possible that these effects are being driven by the large trial by Hassanian-Moghaddam and colleagues that tested a brief contact intervention in Iran [68]. This finding is in contrast to a review by Ougrin and colleagues, which found evidence of benefit for clinical interventions in reducing the proportion of adolescents re-engaging in repeat self-harm [143]. This variation in findings may be explained by the settings in which the studies were conducted, or may be attributable to methodological differences such as the more specific inclusion criteria employed by the current review and/or differences in reporting of results (i.e., use of relative vs absolute effect size). There was also strong evidence of a small effect on suicidal ideation at post-intervention, and to a lesser extent at follow-up, again possibly being driven by the large Hassanian-Moghaddam trial [68].

There is less evidence for interventions delivered in educational or workplace settings given that fewer methodologically-rigorous studies have been conducted. Of note are the large studies conducted by Wasserman and colleagues [101] and Schilling and colleagues [99]. The educational components of the interventions tested in these studies appeared to reduce self-harm at post-intervention and at follow-up [99–101], although there were too few studies to conduct meaningful sub-group analyses. There was also an effect on suicidal ideation at post-intervention, but not follow-up. Overall these results indicate that school-based psycho-educational interventions that are coupled with screening have the potential to be effective, however the robustness of findings is hampered by study quality.

To some extent the overall limited effects detected may reflect a lack of statistical power, either due to small sample sizes at baseline or high attrition rates. Many studies (in particular those of indicated interventions) were underpowered and did not find statistically significant improvements despite the direction of effect being positive. This was particularly true for studies examining self-harm given the large sample sizes required to detect an effect [144]. It may also be that suicidal ideation and self-harm are different constructs, and whilst it is largely accepted that they exist along a continuum [145], specific processes may facilitate the transition from suicidal ideation to suicide attempt [146]. It may therefore be the case that existing interventions more effectively target suicidal ideation than self-harm, and that interventions with

stronger theoretical underpinnings are required to reduce self-harm and suicide. Further work delineating the modifiable risk and protective factors associated with repeated self-harm is therefore required [147].

Evidence regarding the efficacy of interventions in community settings was mixed. The studies that examined the impact of multimodal interventions generally reported reductions in rates of suicide and/or self-harm, although study quality was variable. These findings are encouraging given the emphasis in many countries on place-based responses to suicide prevention [148,149]. The interventions tested typically comprised universal educational programs, gatekeeper training, screening, and treatment responses where appropriate, and appeared to positively impact young people. These intervention types should be included in future place-based approaches and subject to rigorous testing.

Means restriction, such as reducing access to known jumping sites, has long been considered an effective suicide prevention intervention [17,18]. Our review identified few studies examining the effects of means restriction on young people, and those that did focused on firearm restriction. These were generally associated with decreases in rates of firearm suicide, but no reduction in overall youth suicides. An explanation may be that firearm suicides are relatively uncommon among youth in the countries studied. For example, three studies were conducted in Canada where the most common method of youth suicide is hanging [150]. It stands to reason that restricting access to a particular method will only reduce overall suicide rates if it is a method commonly used by the population.

Despite the spread of studies across intervention types and settings, gaps existed. For example, General Practitioners (GPs) are often a first port of call for young people yet there were no studies in primary care settings. GPs have identified the need for training in youth suicide prevention [151]; as such primary care settings may provide an opportunity for intervention early in the suicidal trajectory that is currently being missed. Additionally, few studies were conducted in universities or workplaces compared to schools. Given that suicide rates are highest post-school age [152], tertiary education facilities and workplaces are key settings for future suicide prevention efforts and greater evidence is required [142,153]. Moreover, only six studies tested online interventions; all were in educational settings. There is increasing evidence supporting the efficacy of online interventions in the treatment of depression and anxiety [154], as well as evidence supporting their acceptability with young people at risk of suicide and potential to reduce risk [155]. All the studies of online interventions were CBT-based and most appeared to show promise, raising the question of why online interventions are not being trialed in clinical settings. This is an important avenue for youth suicide prevention yet to be capitalized on.

Finally, there are some groups who are underrepresented in this research. Only three studies [93,94,138] tested interventions among indigenous young people, despite this group being at elevated risk in many countries [156]. Similarly, same-sex-attracted and gender diverse young people are at elevated risk of suicide [157], yet only one study specifically targeted same-sex attracted youth [83]. Whilst this may be partially due to methodological challenges [156,158], generating evidence regarding effective suicide prevention approaches for these populations must be a priority. Related to this, females were over-represented in the studies reviewed. This is unsurprising given the higher rates of both self-harm and help-seeking among females compared to males [159,160], however there is a lack of knowledge regarding effective interventions for young men, whose rates of suicide are three times those of females [1].

A strength of this review is the inclusion criteria used. These were both broad (e.g., no restrictions on intervention approach or study design) and specific (i.e., studies tested interventions that were specifically designed for suicide prevention and reported suicide-related outcome data). Whilst some potentially effective interventions may have been excluded (e.g., those designed to treat or prevent depression), this review is well-placed to provide guidance regarding what

does and does not impact suicide-related outcomes in young people. Despite this, some limitations must be addressed.

Firstly, the broad scope of the review, together with time and resource constraints, required us to make a number of pragmatic methodological decisions. For example, we adopted a pragmatic approach to assessing study quality, as applying standard Risk of Bias criteria to the non-RCTs would result in a low quality rating for all studies. Although we acknowledge the high risk of bias associated with non-randomized study designs, ethical and methodological barriers often prevent suicide prevention researchers from conducting RCTs. To accommodate this, the quality of non-RCTs was assessed using a tool appropriate to that design. Overall, however, study quality was limited. Indeed, many RCTs were not reported according to the Consort statement [161] and many were underpowered. Whilst this is not uncommon in suicide prevention research [144], priority needs to be given to well-designed, sufficiently powered studies. Additionally, for pragmatic reasons we did not include analysis of publication bias in our analysis of study quality. Other minor methodological limitations relate to our decisions not to prospectively register the review and not to contact key authors in the field. Although these steps are encouraged, they are not a requirement of compliance with the PRISMA statement and were not anticipated to impact the results; therefore due to time and resource constraints they were not a part of the present review.

A third limitation relates to the quality of the studies included in the meta-analysis, the results of which should be treated with caution. Additionally, on several occasions different studies contributed data to the post-intervention and follow-up outcomes. We therefore cannot be certain that changes at follow-up are in fact the result of a true reduction in the treatment effect over time. There was also heterogeneity in the control conditions and in the outcome measures used between studies, limiting our ability to be confident that studies measured the same constructs. For example, methods to assess self-harm included self-report instruments, hospital data and clinician-rated interviews. It was also often unclear if measures had been validated among young people. Researchers have previously called for the use of well-validated and standardized measures in adult suicide research, and we argue the same is required in studies with youth [162].

Finally, we acknowledge that a number of relevant studies have been published since the search was conducted. For example, a 2018 RCT trial found no benefit of systemic family therapy compared to treatment as usual in reducing subsequent hospital presentations for young people who self-harm [163]. Another RCT found DBT was more effective in reducing repeat suicide attempts in adolescents, compared to individual and group supportive therapy [164]. Although these studies both meet criteria for inclusion in the current review they were published after our search was conducted.

5. Conclusion

This review identified a large number of studies testing a broad range of interventions across multiple settings. We found that some interventions for example, brief contact interventions in clinical settings, and psychoeducation combined with screening in school settings can reduce the frequency of self-harm and suicidal ideation, although it is likely the size of these studies that is driving the effects. Large-scale multimodal interventions also show promise. Despite these promising findings there remains a paucity of high-quality youth suicide prevention intervention studies. Whilst not all interventions lend themselves to testing via RCTs, other robust study designs can and should be employed. Additionally, many studies, particularly those in clinical and community settings, tend to test interventions originally designed for adults. By focusing suicide prevention efforts on generic, as opposed to youth-specific, interventions, we are likely missing crucial opportunities for intervention, such as delivery via online platforms. Future research should adapt known effective interventions for young people,

and for delivery online. A focus on university and workplace settings is also warranted.

Although young people have repeatedly been identified by suicide prevention policy as a group requiring specific attention, their suicide rates are rising. To reverse this trend, we need more large-scale methodologically-rigorous studies that develop and test new approaches. These approaches should be acceptable to *all* young people and capitalize on the ways in which young people interact with the health system, supports, and services.

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Outstanding Questions

- Despite the encouraging findings, key questions remain as to exactly which components of interventions, in particular those delivered in clinical settings, are most effective when it comes to reducing suicide risk among young people.
- There is also a pressing need for large-scale high quality trials in clinical, educational and community settings. This includes in primary care, tertiary education and online settings, which are currently largely neglected.
- Questions also remain as to what interventions are most likely to be effective in sub-sections of the population, including among indigenous young people, those who live in low to middle income countries, and those who identify as same sex attracted and/or gender diverse.

Author Contributions

Jo Robinson obtained funds for the study. She oversaw the design and conduct of the review, including data extraction, analysis and interpretation. She wrote the manuscript. She also played a leading role in the design of the search strategy.

Eleanor Bailey was responsible for conducting the literature search and coordinating the screening and data extraction phases. She also assisted with designing the search strategy, screening, data extraction, interpretation of results and preparation of the manuscript.

Katrina Witt assisted with screening and data extraction, and was responsible for conducting and interpreting the meta-analysis together with Sarah Hetrick.

Nina Stefanac assisted with screening, data extraction and preparation of the manuscript.

Allison Milner assisted with screening and data extraction.

Dianne Currier assisted with screening and data extraction.

Jane Pirkis provided methodological and conceptual advice. She also contributed to writing the manuscript.

Patrick Condon assisted with the development of the search strategy.

Sarah Hetrick assisted with the development of the search, assisted with screening and data extraction, was responsible for conducting

and interpreting the meta-analysis together with Katrina Witt, and provided general oversight to the study.

References

- [1] World Health Organization. Preventing suicide: a global imperative. Switzerland: World Health Organization; 2014.
- [2] Hawton K, Zahl D, Weatherall R. Suicide following deliberate self-harm: long-term follow-up of patients who presented to a general hospital. *Br J Psychiatry* 2003;182(6):537–42.
- [3] Geulayov G, Kapur N, Turnbull P, et al. Epidemiology and trends in non-fatal self-harm in three centres in England, 2000–2012: findings from the Multicentre Study of Self-harm in England. *BMJ Open* 2016;6(4).
- [4] Nock M, Green J, Hwang I, et al. Prevalence, correlates, and treatment of lifetime suicidal behavior among adolescents: results from the national comorbidity survey replication adolescent supplement. *JAMA Psychiat* 2013;70(3):300–10.
- [5] Hawton K, Harriss L. Deliberate self-harm in young people: characteristics and subsequent mortality in a 20-year cohort of patients presenting to hospital. *J Clin Psychiatry* 2007;68:1574–83.
- [6] Hunt IM, Kapur N, Webb R, et al. Suicide in recently discharged psychiatric patients: a case-control study. *Psychol Med* 2009;39(3):443–9.
- [7] Commonwealth Government Department of Health and Aging. Living is for everyone (LiFE) framework. Canberra, Australia: Author; 2007.
- [8] New Zealand Government. New Zealand suicide prevention action plan 2013–2016. New Zealand: Author; 2013.
- [9] U.S. Department of Health and Human Services. National Strategy for Suicide Prevention: goals and objectives for action. Washington, D.C.: Office of the Surgeon General and National Action Alliance for Suicide Prevention; 2012.
- [10] Patton GC, Sawyer SM, Santelli JS, et al. Our future: a Lancet commission on adolescent health and wellbeing. *Lancet* 2016;387(10036):2423–78.
- [11] National Mental Health Commission. The 2017 report on mental health and suicide prevention. Author: Sydney, Australia; 2017.
- [12] Hawton K, Pirkis J. Suicide is a complex problem that requires a range of prevention initiatives and methods of evaluation. *Br J Psychiatry* 2017;210(6):381.
- [13] Goldney R. Suicide prevention: a pragmatic review of recent studies. *Crisis* 2005;26:128–40.
- [14] Callear A, Christensen H, Freeman A, et al. A systematic review of psychosocial suicide prevention interventions for youth. *Eur Child Adolesc Psychiatry* 2016;25:467–82.
- [15] Robinson J, Cox G, Malone A, et al. A systematic review of school based interventions aimed at preventing, treating, and responding to, suicide-related behaviour in young people. *Crisis* 2012;28:1–19.
- [16] Isaac M, Elias B, Katz L, et al. Gatekeeper training as a preventative intervention for suicide: a systematic review. *Can J Psychiatr* 2009;54(4):260–8.
- [17] Mann J, Apter A, Bertolote J, et al. Suicide prevention strategies: a systematic review. *JAMA* 2005;294:2064–74.
- [18] Zalsman G, Hawton K, Wasserman D, et al. Suicide prevention strategies revisited: 10-year systematic review. *Lancet Psychiatry* 2017;3:646–59.
- [19] Higgins J, Altman D, Sterne J. Assessing risk of bias in included studies. In: Higgins J, editor. *Cochrane handbook for systematic reviews of interventions*. The Cochrane Collaboration; 2011.
- [20] Moher D, Liberati A, Tetzlaff J, Altman D. Preferred reporting items for systematic reviews and meta-analyses: The PRISMA Statement. *PLoS Med* 2009;6(6):37–46.
- [21] Cohen J. A coefficient of agreement for nominal scales. *Educ Psychol Meas* 1960;20:37–46.
- [22] Mak H, Yau K, Chan B. Prevalence-adjusted bias-adjusted kappa values as additional indicators to measure observer agreement. *Radiology* 2004;232:302–3.
- [23] Robinson J, Hetrick S, Martin C. Preventing suicide in young people: systematic review. *Aust N Z J Psychiatry* 2011;45(1):3–26.
- [24] Hawton K, Witt KG, Taylor Salisbury TL, et al. Interventions for self-harm in children and adolescents. *Cochrane Database Syst Rev* 2015;12.
- [25] Norwegian Knowledge Centre for the Health Services. Effective Practice and Organisation of Care (EPOC). EPOC Resources for review authors Oslo; 2015.
- [26] Cochrane Effective Practice and Organisation of Care (EPOC). Suggested risk of bias criteria for EPOC reviews. <http://epoc.cochrane.org/resources/epoc-resources-review-authors>; 2017.
- [27] Hawton K, Bergen H, Cooper J, et al. Suicide following self-harm: Findings from the Multicentre Study of self-harm in England, 2000–2012. *J Affect Disord* 2015;175:147–51.
- [28] Rücker G, Cates CJ, Schwarzer G. Methods for including information from multi-arm trials in pairwise meta-analysis. *Res Synth Methods* 2017;8(4):392–403.
- [29] Pace NL. Research methods for meta-analyses. *Clinical Anaesthesiology* 2011;25(4):523–33.
- [30] Dersimonian R, Laird N. Meta-analysis in clinical trials. *Control Clin Trials* 1986;7(1):177–88.
- [31] Borenstein M, Hedges LV, Higgins JPT, Rothstein HR. Comprehensive meta-analysis. 2.2.064 ed. Englewood, NJ: Biostat, Inc.; 2011.
- [32] Higgins JP, Thompson SG, Deeks JJ, Altman DG. Measuring inconsistency in meta-analyses. *Br Med J* 2003;327:557–60.
- [33] Milner A, Carter G, Pirkis J, Robinson J, Spittal M. Letters, green cards, telephone calls, and postcards: a systematic and meta-analytic review of brief contact interventions for reducing self-harm, suicide attempts, and suicide. *Br J Psychiatry* 2015;206:184–90.
- [34] Fleischmann A, Bertolote JM, Wasserman D, et al. Effectiveness of brief intervention and contact for suicide attempters: a randomized controlled trial in five countries. *Bull World Health Organ* 2008;86(9):703–9.

- [35] Rotheram-Borus MJ, Piacentini J, Cantwell C, Belin TR, Song J. The 18-month impact of an emergency room intervention for adolescent female suicide attempters. *J Consult Clin Psychol* 2000;68(6):1081–93.
- [36] Eggert L, Thompson E, Herting J, Nicholas L. Reducing suicide potential among high-risk youth: tests of a school-based prevention program. Erratum. *Suicide Life Threat Behav* 1999;29(96).
- [37] Silverstone PH, Bercov M, Suen VYM, et al. Long-term results from the Empowering a Multimodal Pathway Toward Healthy Youth Program, a multimodal school-based approach, show marked reductions in suicidality, depression, and anxiety in 6227 students in grades 6–12 (aged 11–18). *Front Psychiatry* 2017;8:81.
- [38] Walrath C, Garraza LG, Reid H, Goldston DB, McKeon R. Impact of the Garrett Lee Smith youth suicide prevention program on suicide mortality. *Am J Public Health* 2015;105(5):986–93.
- [39] Ougrin D, Boege I, Stahl D, Banarsee R, Taylor E. Randomised controlled trial of therapeutic assessment versus usual assessment in adolescents with self-harm: 2-year follow-up. *Arch Dis Child* 2013;98(10):772–6.
- [40] Rudd MD, Rajab MH, Orman DT, Joiner T, Stulman DA, Dixon W. Effectiveness of an outpatient intervention targeting suicidal young adults: preliminary results. *J Consult Clin Psychol* 1996;64(1):179–90.
- [41] Alavi A, Sharifi B, Ghanizadeh A, Dehbozorgi G. Effectiveness of cognitive-behavioral therapy in decreasing suicidal ideation and hopelessness of the adolescents with previous suicidal attempts. *Iran J Pediatr* 2013;23(4):467–72.
- [42] Asarnow JR, Hughes JL, Babeva KN, Sugar CA. Cognitive-behavioral family treatment for suicide attempt prevention: a randomized controlled trial. *J Am Acad Child Adolesc Psychiatry* 2017;56(6):506–14.
- [43] Byford S, Harrington R, Torgerson D, et al. Cost-effectiveness analysis of a home-based social work intervention for children and adolescents who have deliberately poisoned themselves. Results of a randomised controlled trial. *Br J Psychiatry* 1999; 174:56–62.
- [44] Carter GL, Willcox CH, Lewin TJ, Conrad AM, Bendit N. Hunter DBT project: randomized controlled trial of dialectical behaviour therapy in women with borderline personality disorder. *Aust N Z J Psychiatry* 2010;44(2):162–73.
- [45] Cooney E, Davis K, Thompson P, Wharewera-Mika J, Stewart J. Feasibility of evaluating DBT for self-harming adolescents: A small randomised controlled trial. Auckland, New Zealand: Te Pou o Te Whakaaro Nui; 2010.
- [46] Diamond GS, Wintersteen MB, Brown GK, et al. Attachment-based family therapy for adolescents with suicidal ideation: a randomized controlled trial. *J Am Acad Child Adolesc Psychiatry* 2010;49(2):122–31.
- [47] Donaldson D, Spirito A, Esposito-Smythers C. Treatment for adolescents following a suicide attempt: results of a pilot trial. *J Am Acad Child Adolesc Psychiatry* 2005;44 (2):113–20.
- [48] Esposito-Smythers C, Spirito A, Kahler CW, Hunt J, Monti P. Treatment of co-occurring substance abuse and suicidality among adolescents: a randomized trial. *J Consult Clin Psychol* 2011;79(6):728–39.
- [49] Green JM, Wood AJ, Kerfoot MJ, et al. Group therapy for adolescents with repeated self harm: randomised controlled trial with economic evaluation. *BMJ* 2011;342: d682.
- [50] Harrington R, Kerfoot M, Dyer E, et al. Randomized trial of a home-based family intervention for children who have deliberately poisoned themselves. *J Am Acad Child Adolesc Psychiatry* 1998;37(5):512–8.
- [51] Hazell PL, Martin G, McGill K, et al. Group therapy for repeated deliberate self-harm in adolescents: failure of replication of a randomized trial. *J Am Acad Child Adolesc Psychiatry* 2009;48(6):662–70.
- [52] Huey Jr SJ, Henggeler SW, Rowland MD, et al. Multisystemic therapy effects on attempted suicide by youths presenting psychiatric emergencies. *J Am Acad Child Adolesc Psychiatry* 2004;43(2):183–90.
- [53] Husain N, Afsar S, Ara J, et al. Brief psychological intervention after self-harm: randomised controlled trial from Pakistan. *Br J Psychiatry* 2014;204(6):462–70.
- [54] King CA, Kramer A, Preuss L, Kerr DC, Weiss L, Venkataraman S. Youth-Nominated Support Team for Suicidal Adolescents (version 1): a randomized controlled trial. *J Consult Clin Psychol* 2006;74(1):199–206.
- [55] King CA, Klaus N, Kramer A, Venkataraman S, Quinlan P, Gillespie B. The Youth-Nominated Support Team-Version II for suicidal adolescents: a randomized controlled intervention trial. *J Consult Clin Psychol* 2009;77(5):880–93.
- [56] King CA, Gipson PY, Horwitz AG, Opperman KJ. Teen Options for Change (TOC): an intervention for adolescent emergency patients who screen positive for suicide risk. *Psychiatr Serv* 2015;66(1):97–100.
- [57] McLeavey BC, Daly RJ, Ludgate JW, Murray CM. Interpersonal problem-solving skills training in the treatment of self-poisoning patients. *Suicide Life Threat Behav* 1994;24(4):382–94.
- [58] Mehlum L, Ramberg M, Tormoen AJ, et al. Dialectical behavior therapy compared with enhanced usual care for adolescents with repeated suicidal and self-harming behavior: outcomes over a one-year follow-up. *J Am Acad Child Adolesc Psychiatry* 2016;55(4):295–300.
- [59] Pineda J, Dadds MR. Family intervention for adolescents with suicidal behavior: a randomized controlled trial and mediation analysis. *J Am Acad Child Adolesc Psychiatry* 2013;52(8):851–62.
- [60] Power PJ, Bell RJ, Mills R, et al. Suicide prevention in first episode psychosis: the development of a randomised controlled trial of cognitive therapy for acutely suicidal patients with early psychosis. *Aust N Z J Psychiatry* 2003;37(4):414–20.
- [61] Rossouw TI, Fonagy P. Mentalization-based treatment for self-harm in adolescents: a randomized controlled trial. *J Am Acad Child Adolesc Psychiatry* 2012;51(12) [1304–13.e3].
- [62] Slee N, Garnefski N, van der Leeden R, Arensman E, Spinhoven P. Cognitive-behavioural intervention for self-harm: randomised controlled trial. *Br J Psychiatry* 2008;192(3):202–11.
- [63] Wharff EA, Ginnis KB, Ross AM, White EM, White MT, Forbes PW. Family-based crisis intervention with suicidal adolescents: a randomized clinical trial. *Pediatr Emerg Care* 2017. <https://doi.org/10.1097/PEC.0000000000001076> [Epub ahead of print].
- [64] Wood A, Trainor G, Rothwell J, Moore A, Harrington R. Randomized trial of group therapy for repeated deliberate self-harm in adolescents. *J Am Acad Child Adolesc Psychiatry* 2001;40(11):1246–53.
- [65] Asarnow JR, Baraff LJ, Berk M, et al. An emergency department intervention for linking pediatric suicidal patients to follow-up mental health treatment. *Psychiatr Serv* 2011;62(11):1303–9.
- [66] Bertolote JM, Fleischmann A, De Leo D, et al. Repetition of suicide attempts: data from emergency care settings in five culturally different low- and middle-income countries participating in the WHO SUPRE-MISS study. *Crisis* 2010;31(4):194–201.
- [67] Cotgrove A, Zirinsky L, Black D, Weston D. Secondary prevention of attempted suicide in adolescence. *J Adolesc* 1995;18(5):569–77.
- [68] Hassanian-Moghaddam H, Sarjani S, Kolahi AA, Carter GL. Postcards in Persia: randomised controlled trial to reduce suicidal behaviours 12 months after hospital-treated self-poisoning. *Br J Psychiatry* 2011;198(4):309–16.
- [69] Ougrin D, Zundel T, Banarsee R, Bottle A, Taylor E. Trial of Therapeutic Assessment in London: randomised controlled trial of Therapeutic Assessment versus standard psychosocial assessment in adolescents presenting with self-harm. *Arch Dis Child* 2011;96(2):148–53.
- [70] Robinson J, Yuen H, Gook S, et al. Can receipt of a regular postcard reduce suicide-related behaviour in young help seekers? A randomized controlled trial. *Early Interv Psychiatry* 2012;6(2):145–52.
- [71] Spirito A, Boergers J, Donaldson D, Bishop D, Lewander W. An intervention trial to improve adherence to community treatment by adolescents after a suicide attempt. *J Am Acad Child Adolesc Psychiatry* 2002;41(4):435–42.
- [72] Spirito A, Wolff JC, Seaboyer LM, et al. Concurrent treatment for adolescent and parent depressed mood and suicidality: feasibility, acceptability, and preliminary findings. *J Child Adolesc Psychopharmacol* 2015;25(2):131–9.
- [73] Asarnow JR, Berk M, Hughes JL, Anderson NL. The SAFETY Program: a treatment-development trial of a cognitive-behavioral family treatment for adolescent suicide attempters. *J Clin Child Adolesc Psychol* 2015;44(1):194–203.
- [74] Courtney DB, Flament MF. Adapted dialectical behavior therapy for adolescents with self-injurious thoughts and behaviors. *J Nerv Ment Dis* 2015;203(7):537–44.
- [75] Duarte-Velez Y, Torres-Davila P, Spirito A, Polanco N, Bernal G. Development of a treatment protocol for Puerto Rican adolescents with suicidal behaviors. *Psychotherapy* 2016;53(1):45–56.
- [76] Esposito-Smythers C, Spirito A, Uth R, Lachance H. Cognitive behavioral treatment for suicidal alcohol abusing adolescents: development and pilot testing. *Am J Addict* 2006;15(Suppl. 1):126–30.
- [77] Geddes K, Dziurawiec S, Lee CW. Dialectical behaviour therapy for the treatment of emotion dysregulation and trauma symptoms in self-injurious and suicidal adolescent females: a pilot programme within a community-based child and adolescent mental health service. *Psychiatry* 2013;2013:10.
- [78] Gutstein SE, Rudd MD. An outpatient treatment alternative for suicidal youth. *J Adolesc* 1990;13(3):265–77.
- [79] James AC, Winnill L, Anderson C, Alfoadari K. A preliminary study of an extension of a community dialectic behaviour therapy (DBT) programme to adolescents in the looked after care system. *Child Adolesc Mental Health* 2011;16(1):9–13.
- [80] James S, Freeman KR, Mayo D, et al. Does insurance matter? Implementing dialectical behavior therapy with two groups of youth engaged in deliberate self-harm. *Admin Pol Ment Health* 2015;42(4):449–61.
- [81] King R, Nurcombe B, Bickman L, Hides L, Reid W. Telephone counselling for adolescent suicide prevention: changes in suicidality and mental state from beginning to end of a counselling session. *Suicide Life Threat Behav* 2003;33(4):400–11.
- [82] Cwik MF, Tingey L, Lee A, et al. Development and piloting of a brief intervention for suicidal American Indian adolescents. *Am Indian Alsk Native Ment Health Res* 2016;23(1):105–24.
- [83] Diamond GM, Diamond GS, Levy S, Closs C, Ladipo T, Siqueland L. Attachment-based family therapy for suicidal lesbian, gay, and bisexual adolescents: a treatment development study and open trial with preliminary findings. *Psychol Sex Orientat Gend Divers* 2013;1(5):91–100.
- [84] Katz LY, Cox BJ, Gunasekara S, Miller AL. Feasibility of dialectical behavior therapy for suicidal adolescent inpatients. *J Am Acad Child Adolesc Psychiatry* 2004;43(3): 276–82.
- [85] Law YW, Yip PS, Lai CC, et al. A pilot study on the efficacy of volunteer mentorship for young adults with self-harm behaviors using a quasi-experimental design. *Crisis* 2016;37(6):415–26.
- [86] Oldershaw A, Simic M, Grima E, et al. The effect of cognitive behavior therapy on decision making in adolescents who self-harm: a pilot study. *Suicide Life Threat Behav* 2012;42(3):255–65.
- [87] Perera EA, Kathirarachchi ST. Problem-solving counseling as a therapeutic tool on youth suicidal behavior in the suburban population in Sri Lanka. *Indian J Psychiatry* 2011;53(1):30–5.
- [88] Rathus JH, Miller AL. Dialectical behavior therapy adapted for suicidal adolescents. *Suicide Life Threat Behav* 2002;32(2):146–57.
- [89] Rotheram-Borus MJ, Piacentini J, Van Rossem R, et al. Enhancing treatment adherence with a specialized emergency room program for adolescent suicide attempters. *J Am Acad Child Adolesc Psychiatry* 1996;35(5):654–63.
- [90] Wharff EA, Ginnis KM, Ross AM. Family-based crisis intervention with suicidal adolescents in the emergency room: a pilot study. *Soc Work* 2012;57(2):133–43.
- [91] Brent DA, Greenhill LL, Compton S, et al. The Treatment of Adolescent Suicide Attempters study (TASA): predictors of suicidal events in an open treatment trial. *J Am Acad Child Adolesc Psychiatry* 2009;48(10):987–96.

- [92] Orbach I, Bar-Joseph H. The impact of a suicide prevention program for adolescents on suicidal tendencies, hopelessness, ego identity, and coping. *Suicide Life Threat Behav* 1993;23(2):120–9.
- [93] Lafromboise T, Howard-Pitney B. The Zuni Life Skills Development Curriculum: a collaborative approach to curriculum development. *Am Indian Alsk Native Ment Health Res Monogr Ser* 1994;4:98–121.
- [94] Lafromboise T, Howard-Pitney B. The Zuni Life Skills Development Curriculum: description and evaluation of a suicide prevention program. *J Couns Psychol* 1995;42(4):479–86.
- [95] Eggert LL, Thompson EA, Randell BP, Pike KC. Preliminary effects of brief school-based prevention approaches for reducing youth suicide-risk behaviors, depression, and drug involvement. *J Child Adolesc Psychiatr Nurs* 2002;15(2):48–64.
- [96] Hooven C, Walsh E, Pike KC, Herting JR. Promoting CARE: including parents in youth suicide prevention. *Fam Community Health* 2012;35(3):225–35.
- [97] Robinson WL, Case MH, Whipple CR, et al. Culturally grounded stress reduction and suicide prevention for African American adolescents. *Pract Innov (Wash DC)* 2016;1(2):117–28.
- [98] Tang TC, Jou SH, Ko CH, Huang SY, Yen CF. Randomized study of school-based intensive interpersonal psychotherapy for depressed adolescents with suicidal risk and parasuicide behaviors. *Psychiatry Clin Neurosci* 2009;63(4):463–70.
- [99] Schilling EA, Aseltine Jr RH, James A. The SOS Suicide Prevention Program: further evidence of efficacy and effectiveness. *Prev Sci* 2016;17(2):157–66.
- [100] Schilling EA, Lawless M, Buchanan L, Aseltine Jr RH. "Signs of Suicide" shows promise as a middle school suicide prevention program. *Suicide Life Threat Behav* 2014;44(6):653–67.
- [101] Wasserman D, Hoven CW, Wasserman C, et al. School-based suicide prevention programmes: the SEYLE cluster-randomised, controlled trial. *Lancet* 2015;385(9977):1536–44.
- [102] King KA, Strunk CM, Sorter MT. Preliminary effectiveness of surviving the teens suicide prevention and depression awareness program on adolescents' suicidality and self-efficacy in performing help-seeking behaviors. *J Sch Health* 2011;81(9):581–90.
- [103] Vieland V, Whittle B, Garland A, Hicks R, Shaffer D. The impact of curriculum-based suicide prevention programs for teenagers: an 18-month follow-up. *J Am Acad Child Adolesc Psychiatry* 1991;30(5):811–5.
- [104] Hazell P, Lewin T. An evaluation of postvention following adolescent suicide. *Suicide Life Threat Behav* 1993;23(2):101–9.
- [105] Biddle VS, Kern 3rd J, Brent DA, Thurekettle MA, Puskas KR, Sekula LK. Student assistance program outcomes for students at risk for suicide. *J Sch Nurs* 2014;30(3):173–86.
- [106] Eggert LL, Thompson EA, Herting JR, Nicholas LJ. Reducing suicide potential among high-risk youth: tests of a school-based prevention program. *Suicide Life Threat Behav* 1995;25(2):276–96.
- [107] Aseltine Jr RH, James A, Schilling EA, Glanovsky J. Evaluating the SOS suicide prevention program: a replication and extension. *BMC Public Health* 2007;7:161.
- [108] Silverstone PH, Bercov M, Suen VYM, et al. Initial findings from a novel school-based program, EMPATHY, which may help reduce depression and suicidality in youth. *PLoS One* 2015;10(5):e0125527.
- [109] Zenere 3rd FJ, Lazarus PJ. The decline of youth suicidal behavior in an urban, multicultural public school system following the introduction of a suicide prevention and intervention program. *Suicide Life Threat Behav* 1997;27(4):387–402.
- [110] Bailey E, Spittal MJ, Pirkis J, Gould M, Robinson J. Universal suicide prevention in young people: an evaluation of the safeTALK program in Australian high schools. *Crisis* 2017;38(5):300–8.
- [111] Hetrick S, Yuen HP, Cox G, et al. Does cognitive behavioural therapy have a role in improving problem solving and coping in adolescents with suicidal ideation? *Cogn Behav Ther* 2014;7.
- [112] Robinson J, Hetrick S, Cox G, et al. Can an internet-based intervention reduce suicidal ideation, depression and hopelessness among secondary school students: results from a pilot study. *Early Interv Psychiatry* 2016;10(1):28–35.
- [113] Guille C, Zhao Z, Krystal J, Nichols B, Brady K, Sen S. Web-based cognitive behavioral therapy intervention for the prevention of suicidal ideation in medical interns: a randomized clinical trial. *JAMA Psychiat* 2015;72(12):1192–8.
- [114] Kovac SH, Range LM. Does writing about suicidal thoughts and feelings reduce them? *Suicide Life Threat Behav* 2002;32(4):428–40.
- [115] Fitzpatrick KK, Witte TK, Schmidt NB. Randomized controlled trial of a brief problem-orientation intervention for suicidal ideation. *Behav Ther* 2005;36(4):323–33.
- [116] Pistorello J, Fruzzetti AE, MacLaine C, Gallop R, Iverson KM. Dialectical behavior therapy (DBT) applied to college students: a randomized clinical trial. *J Consult Clin Psychol* 2012;80(6):982–94.
- [117] Lerner MS, Clum GA. Treatment of suicide ideators: a problem-solving approach. *Behav Ther* 1990;21(4):403–11.
- [118] Joffe P. An empirically supported program to prevent suicide in a college student population. *Suicide Life Threat Behav* 2008;38(1):87–103.
- [119] Till B, Tran US, Voracek M, Niederkrotenthaler T. Beneficial and harmful effects of educational suicide prevention websites: randomised controlled trial exploring Papageno v. Werther effects. *Br J Psychiatry* 2017;211(2):109–15.
- [120] Shalef L, Tatsa-Laur L, Derazne E, Mann JJ, Fruchter E. An effective suicide prevention program in the Israeli Defense Forces: a cohort study. *Eur Psychiatry* 2016;31:37–43.
- [121] McDaniel WW, Rock M, Grigg JR. Suicide prevention at a United States Navy training command. *Mil Med* 1990;155(4):173–5.
- [122] Hill RM, Pettit JW. Pilot randomized controlled trial of LEAP: a selective preventive intervention to reduce adolescents' perceived burdensomeness. *J Clin Child Adolesc Psychol* 2016;1–12.
- [123] Hetrick SE, Yuen HP, Bailey E, et al. Internet-based cognitive behavioural therapy for young people with suicide-related behaviour (Reframe-IT): a randomised controlled trial. *Evid Based Ment Health* 2017;20(3):76–82.
- [124] Ahmadi A, Ytterstad B. Prevention of self-immolation by community-based intervention. *Burns* 2007;33(8):1032–40.
- [125] Beautrais AL, Fergusson DM, Horwood LJ. Firearms legislation and reductions in firearm-related suicide deaths in New Zealand. *Aust N Z J Psychiatry* 2006;40(3):253–9.
- [126] Caron J. Gun control and suicide: possible impact of Canadian legislation to ensure safe storage of firearms. *Arch Suicide Res* 2004;8(4):361–74.
- [127] Center for Disease Control. Suicide prevention evaluation in a Western Athabaskan American Indian Tribe-New Mexico, 1988–1997. *MMWR Morb Mortal Wkly Rep* 1998;47(13):257–61.
- [128] Cheung AH, Dewa CS. Current trends in youth suicide and firearms regulations. *Can J Public Health* 2005;96(2):131–5.
- [129] Cwik MF, Tingey L, Maschino A, et al. Decreases in suicide deaths and attempts linked to the White Mountain apache suicide surveillance and prevention system, 2001–2012. *Am J Public Health* 2016;106(12):2183–9.
- [130] Hacker K, Collins J, Gross-Young L, Almeida S, Burke N. Coping with youth suicide and overdose: one community's efforts to investigate, intervene, and prevent suicide contagion. *Crisis* 2008;29(2):86–95.
- [131] Leenaars AA, Lester D. The impact of gun control on suicide and homicide across the life span. *Can J Behav Sci* 1997;29(1):1–6.
- [132] Lubin C, Werbeloff N, Halperin D, Shmushkevitch M, Weiser M, Knobler HY. Decrease in suicide rates after a change of policy reducing access to firearms in adolescents: a naturalistic epidemiological study. *Suicide Life Threat Behav* 2010;40(5):421–4.
- [133] Matsubayashi T, Ueda M. The effect of national suicide prevention programs on suicide rates in 21 OECD nations. *Soc Sci Med* 2011;73(9):1395–400.
- [134] May PA, Serna P, Hurt L, Debruyn LM. Outcome evaluation of a public health approach to suicide prevention in an American Indian tribal nation. *Am J Public Health* 2005;95(7):1238–44.
- [135] Niederkrotenthaler T, Till B, Herberth A, et al. Can media effects counteract legislation reforms? The case of adolescent firearm suicides in the wake of the Austrian firearm legislation. *J Adolesc Health* 2009;44(1):90–3.
- [136] Wheeler BW, Gunnell D, Metcalfe C, Stephens P, Martin RM. The population impact on incidence of suicide and non-fatal self harm of regulatory action against the use of selective serotonin reuptake inhibitors in under 18s in the United Kingdom: ecological study. *BMJ* 2008;336(7643):542–5.
- [137] Wheeler BW, Metcalfe C, Martin RM, Gunnell D. International impacts of regulatory action to limit antidepressant prescribing on rates of suicide in young people. *Pharmacoepidemiol Drug Saf* 2009;18(7):579–88.
- [138] Allen J, Rasmus SM, Fok CCT, Charles B, Henry D. Multi-level cultural intervention for the prevention of suicide and alcohol use risk with Alaska Native youth: a nonrandomized comparison of treatment intensity. *Prev Sci* 2017. <https://doi.org/10.1007/s11211-017-0798-9> [Epub ahead of print].
- [139] Garraza L, Walrath C, Goldston DB, Reid H, McKeon R. Effect of the garrett lee smith memorial suicide prevention program on suicide attempts among youths. *JAMA Psychiat* 2015;72(11):1143–9.
- [140] Robinson J, Pirkis J. Research priorities in suicide prevention: an examination of Australian-based research 2007–11. *Aust Health Rev* 2014;38(1):18–24.
- [141] Dahl R, Allen N, Ballonoff Suleimna A. Importance of investing in adolescence from a developmental science perspective. *Nature* 2018;554:441–50.
- [142] Robinson J, Bailey E, Browne V, Cox G, Hooper C. Raising the Bar for youth suicide prevention. Melbourne: Orygen: The National Centre of Excellence in Youth Mental Health; 2016.
- [143] Ougrin D, Tranah T, Stahl D, Moran P, Asarnow JR. Therapeutic interventions for suicide attempts and self-harm in adolescents: systematic review and meta-analysis. *J Am Acad Child Adolesc Psychiatry* 2015;54(2) [97–107.e2].
- [144] Arensman E, Townsend E, Hawton K, et al. Psychosocial and pharmacological treatment of patients following deliberate self-harm: the methodological issues involved in evaluating effectiveness. *Suicide Life Threat Behav* 2001;31(2).
- [145] Svetic J, De Leo D. The hypothesis of a continuum in suicidality: a discussion on its validity and practical implications. *Ment Illn* 2012;2:e15.
- [146] O'Connor RC. Towards an integrated motivational-volitional model of suicidal behaviour. In: O'Connor RC, Platt S, Gordon J, editors. *Handbook of suicide prevention: research, policy and practice*. John Wiley & Sons, Ltd.: Chichester, UK; 2011.
- [147] Witt K, Milner A, Spittal M, et al. Population attributable risk of factors associated with the repetition of self-harm behaviour in young people presenting to clinical services: a systematic review and meta-analysis. *Eur Child Adolesc Psychiatry* 2018. <https://doi.org/10.1007/s00787-018-1111-6> [Epub ahead of print].
- [148] Kryszinska K, Batterham P, Tye M, et al. Best strategies for reducing the suicide rate in Australia. *Aust N Z J Psychiatry* 2016;50(2):115–8.
- [149] Hegel U, Rummel-Kluge C, Varnik A, Arensman E, Koburger N. Alliances against depression – a community based approach to target depression and to prevent suicidal behaviour. *Neurosci Biobehav Rev* 2013;37:2404–9.
- [150] Statistics Canada. Suicide rates: An overview; 2017.
- [151] Michail M, Tait L, Churchill D. General practitioners' clinical expertise in managing suicidal young people: implications for continued education. *Prim Health Care Res Dev* 2017;18:419–28.
- [152] Australian Bureau of Statistics. 3303.0 Causes of Death, Australia. Canberra: Commonwealth of Australia; 2015; 2016.
- [153] Milner A, Page K, Spencer-Thomas S, Lamotagne AD. Workplace suicide prevention: a systematic review of published and unpublished activities. *Health Promot Int* 2015;30(1):29–37.
- [154] Välimäki M, Anttila K, Anttila M, Lahti M. Web-based interventions supporting adolescents and young people with depressive symptoms: systematic review and meta-analysis. *JMIR Mhealth Uhealth* 2017;5(12) [e180-e].
- [155] Witt KG, Spittal MJ, Carter G, et al. Effectiveness of online and mobile telephone applications ("apps") for the self-management of suicidal ideation and

- self-harm: a systematic review and meta-analysis. *BMC Psychiatry* 2017;17 (1):297.
- [156] Clifford AC, Doran CM, Tsey K. A systematic review of suicide prevention interventions targeting indigenous peoples in Australia, United States, Canada and New Zealand. *BMC Public Health* 2013;13(1):463.
 - [157] Turban JLED. Research review: gender identity in youth: treatment paradigms and controversies. *J Child Psychol Psychiatry* 2017. <https://doi.org/10.1111/jcpp.12833> [Epub ahead of print].
 - [158] Silenzio VMB, Duberstein PR, Tang W, Lu N, Tu X, Homan CM. Connecting the invisible dots: reaching lesbian, gay, and bisexual adolescents and young adults at risk for suicide through online social networks. *Soc Sci Med* 2009;69(3):469–74.
 - [159] Wendt D, Shafer K. Gender and attitudes about mental health help seeking: results from National Data. *Health Soc Work* 2016;41(1) [e20–e8].
 - [160] Morgan C, Webb RT, Carr MJ, et al. Incidence, clinical management, and mortality risk following self harm among children and adolescents: cohort study in primary care. *BMJ* 2017;359.
 - [161] Schulz KF, Altman DG, Moher D. CONSORT 2010 statement: updated guidelines for reporting parallel group randomised trials. *BMJ* 2010;340.
 - [162] Batterham P, Ftanou M, Pirkis J, et al. A systematic review and evaluation of measures for suicidal ideation and behaviors in population-based research. *Psychol Assess* 2015;27(2):501–12.
 - [163] Cottrell DJ, Wright-Hughes A, Collinson M, et al. Effectiveness of systemic family therapy versus treatment as usual for young people after self-harm: a pragmatic, phase 3, multicentre, randomised controlled trial. *The Lancet Psychiatry* 2018;5 (3):203–16.
 - [164] McCauley E, Berk MS, Asarnow JR, et al. Efficacy of dialectical behavior therapy for adolescents at high risk for suicide: a randomized clinical trial. *JAMA Psychiat* 2018;75(8):777–85.