Emergency Care Follow-Up for Children and Adolescents With Suicide Attempts or Ideation

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Abbreviations

PICO participants, interventions, comparators, outcomes
PRISMA Preferred Reporting Items for Systematic Reviews and Meta-Analyses
RCT randomized controlled trial
SR systematic review
Key Messages

- Death by suicide is the second-leading cause of death for young people in Canada. Because 10% to 15% of the people seen in hospital emergency departments for a suicide attempt will repeat the attempt within 12 months following their discharge, it is important to identify what interventions are effective for preventing further self-harm during this time.

- Active follow-up care following an emergency department visit for a suicide attempt is common; however, it is unclear what type (e.g., text message, home visit) or timing (e.g., 24 hours, within 72 hours after emergency department discharge) of interventions is most effective for preventing further self-harm in people younger than 18 years. The objective of this review is to summarize the evidence regarding the clinical effectiveness of active contact and follow-up interventions, and the timing and duration of care, for children and adolescents (younger than 18 years) who present to emergency care for suicide attempts or suicide ideations.

- For children and adolescents presenting to the emergency department for suicide attempts or ideation, telephone-based follow-up care initiated within 1 week of discharge may not affect the number of people who completed the full course of postdischarge treatment, the mean number of sessions attended, or the number of suicide deaths. This finding was based on evidence from 2 systematic reviews, each with 1 primary study relevant to this report. The small sample sizes of these studies (N = 64 and N = 97) and their limited or unclear quality should be considered when interpreting these results.

- None of the relevant primary studies within the systematic reviews reported on mental health outcomes (e.g., depression, social functioning) or harms from the intervention.

- No studies were found that evaluated the clinical effectiveness of other methods of active follow-up care or of different time durations of follow-up care for children and adolescents who present to emergency care for suicide attempts or suicide ideations that met our criteria for this review.

- No evidence-based guidelines were identified that provided recommendations about timing, modality, and which health care professionals should be involved in follow-up care for children and adolescents who present to emergency care for suicide attempts or ideations that met our criteria for this review.

- Guidance documents and guidelines for adults generally recommend that follow-up should occur within 48 hours, particularly for those with safety concerns of subsequent self-harm. These guidance documents also suggest different modalities for follow-up during that time, including telephone calls, visits, and electronic communication. Given the higher risk and potential vulnerability experienced by children and adolescents and the absence of formal clinical guidelines, person-centred follow-up care for all children and adolescents within 48 hours should be considered, similar to adults who present with high concerns.
Context and Policy Issues

In Canada, approximately 12 people die by suicide every day.¹ For those aged between 10 and 19 years, death rates by suicide per 100,000 people are 4.9 in females and 7.4 in males;¹ however, published data may underestimate true numbers due to stigma. In a 2019 Canadian survey of 6,800 adolescents (6,750 cisgender and 50 transgender), transgender adolescents were at increased risk of suicidal ideation (risk ratio = 4.95; 95% confidence interval [CI], 3.63 to 6.75) and having attempted suicide (risk ratio = 7.60; 95% CI, 4.76 to 12.10) compared with cisgender adolescents.² Further, the COVID-19 pandemic has had an impact on the mental health of children and adolescents due to stress, unpredictability, and disruptions (e.g., school, social, family). A 2023 meta-analysis identified 42 studies in 18 countries and reported a 22% increase in emergency department visits for attempted suicide compared with prepandemic rates; the mean age of children presenting to the emergency department for an attempted suicide was 11.7 years.³

For every completed suicide in adolescents, there may be as many as 50 to 100 attempts.⁴ Several organizations, including the WHO, state that a previous suicide attempt is the single most important risk factor for suicide.⁵ Other risk factors include psychiatric disorders, family history of mood disorders, history of physical or sexual abuse, exposure to violence, and biologic factors.⁴ An evidence brief by the Mental Health Commission of Canada reported that 10% to 15% of people seen in a hospital emergency department for a suicide attempt will repeat the attempt within 12 months following discharge.⁶ Up to 70% of individuals who survive a suicide attempt do not attend their first outpatient appointment, so several follow-up and regular contact interventions can be implemented to help ensure the continuity of care for these individuals.⁶ These active follow-up interventions include telephone calls, text messages, emails, letters or postcards, and home visits. The National Institute for Health and Care Excellence (NICE) recommends the following: “If there are ongoing safety concerns for the person after an episode of self-harm, the mental health team, GP, team who carried out the psychosocial assessment or the team responsible for their care should provide initial aftercare within 48 hours of the psychosocial assessment.”⁷ Most recommendations regarding follow-up after an episode of self-harm or suicide attempt are developed using evidence generated from adults.

Several systematic reviews (SRs) have been published that evaluate the effectiveness of active follow-up care; however, they also include adult populations and/or other interventions.⁸⁻¹² It is unclear what are the best interventions (e.g., text message, home visit) and the timing of these interventions (e.g., 24 hours, within 72 hours after emergency department discharge) specific to those younger than 18 years. The objective of this review is to summarize the evidence regarding the clinical effectiveness of early active contact and follow-up interventions, and the timing of initial follow-up care after discharge, for children and adolescents (younger than 18 years) who present to emergency care for suicide attempts or suicide ideations.

Research Questions

1. What is the clinical effectiveness of active contact and follow-up interventions for children and adolescents who present to emergency care for suicide attempts or suicide ideations?
2. What is the clinical effectiveness of different time durations for follow-up care for children and adolescents who present to emergency care for suicide attempts or suicide ideations?

3. What are the evidence-based guidelines about the timing, modality, and health care professionals involved in follow-up care for children and adolescents who present to emergency care for suicide attempts or ideations?

**Methods**

**Literature Search Methods**
An information specialist conducted a literature search on key resources, including MEDLINE, PsycINFO, the Cochrane Database of Systematic Reviews, the International HTA Database, the websites of Canadian and major international health technology agencies, as well as a focused internet search. The search approach was customized to retrieve a limited set of results, balancing comprehensiveness with relevancy. The search strategy comprised both controlled vocabulary, such as the National Library of Medicine’s MeSH (Medical Subject Headings), and keywords. Search concepts were developed based on the elements of the research questions and selection criteria. The main search concepts were suicide, emergency settings, and follow-up methods. The search was completed on June 16, 2023, and limited to English-language documents published since January 1, 2018.

**Selection Criteria and Methods**
One reviewer screened citations and selected studies. In the first level of screening, titles and abstracts were reviewed and potentially relevant articles were retrieved and assessed for inclusion. The final selection of full-text articles was based on the inclusion criteria presented in Table 1. Within the included SRs, when PICO (participants, interventions, comparators, outcomes) elements were not described in sufficient detail to be able to determine if a study was relevant to this report, an additional review of the abstracts was performed.

**Table 1: Selection Criteria**

<table>
<thead>
<tr>
<th>Criteria</th>
<th>Description</th>
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<tbody>
<tr>
<td>Population</td>
<td>Children and adolescents (younger than 18 years) with attempted suicide or suicide ideation, presenting to emergency care (emergency departments) who are not subsequently admitted to the hospital</td>
</tr>
</tbody>
</table>
| Intervention | Q1: Active contact and follow-up care, with initial contact within 1 week of emergency department discharge  
Q2: Time durations from emergency department discharge for initial follow-up care (e.g., 24 hours, 48 hours, 72 hours)  
Q3: Follow-up care |
| Comparator | Q1. Usual care (may include passive contact, general outpatient appointment, or no follow-up)  
Q2. Alternative time duration for follow-up care  
Q3. Not applicable |
### Criteria

<table>
<thead>
<tr>
<th>Criteria</th>
<th>Description</th>
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</thead>
</table>
| Outcomes       | Q1 and Q2. Clinical benefits (e.g., suicide behaviour or ideation, mental health outcomes, treatment engagement, or adherence or attendance) and harms  
Q3. Recommendations about follow-up care interventions or practices (e.g., best practices for timing of care, modality of care, health care professionals) |
| Study designs  | • Health technology assessments  
• Systematic reviews  
• Randomized control trials  
• Nonrandomized studies (with comparators)  
• Evidence-based guidelines |

### Exclusion Criteria

Articles were excluded if they did not meet the selection criteria outlined in Table 1, they were duplicate publications, or were published before 2018. SRs in which all relevant studies were captured in other more recent or more comprehensive SRs were excluded. Primary studies retrieved by the search were excluded if they were captured in 1 or more included SRs. Guidelines with unclear methodology were also excluded.

### Critical Appraisal of Individual Studies

The included publications were critically appraised by 1 reviewer using the following tools as a guide: A Measurement Tool To Assess Systematic Reviews 2 (AMSTAR 2)\(^\text{13}\) for SRs. Summary scores were not calculated for the included studies; rather, the strengths and limitations of each included publication were described narratively.

### Summary of Evidence

#### Quantity of Research Available

A total of 515 citations were identified in the literature search. Following screening of titles and abstracts, 489 citations were excluded and 26 potentially relevant reports from the electronic search were retrieved for full-text review. Fourteen potentially relevant publications were retrieved from the grey literature search for full-text review. Of these potentially relevant articles, 34 were excluded for various reasons; 6 SR publications met the inclusion criteria and are included in this report. Figure 1 presents the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA)\(^\text{14}\) flow chart of the study selection. One included primary study overlapped in the SRs by Witt et al.\(^\text{15}\) and Doupnik et al.\(^\text{8}\) but the population was described as “adolescents hospitalized for suicide attempt” in the SR by Doupnik et al.\(^\text{8}\) (which would be excluded) and “patients presenting to hospital for suicide attempt necessitating medical care in either an emergency department or pediatrics ward of a general children's hospital” in the SR by Witt et al.\(^\text{15}\) (which would be included). Because the SR by Witt et al.\(^\text{15}\) is well-reported and has no major limitations, we included the primary study as reported in this SR.
Additional references of potential interest are provided in Appendix 5. This includes primary studies in adults, reviews with participants who are adults, or participants for whom their age was not clearly defined, and a protocol for a randomized controlled trial (RCT). A brief summary of guidance documents describing follow-up protocols for adults visiting emergency care for suicide attempts or ideation is also presented in Appendix 5, Table 5.

Summary of Study Characteristics
All 6 SRs had broader inclusion criteria than the present review, and 4 SRs did not include any primary studies relevant to this review. For example, the SR by Skopp et al. included studies evaluating the Caring Contacts intervention, but there was no restriction on the population (e.g., adults, patients discharged from psychiatric care), and first contact may have been beyond 1 week after emergency department discharge. This was common among the other SRs, which included broader populations (e.g., adults, patients, hospitalized patients, adolescents presenting to emergency departments for nonpsychiatric reasons), irrelevant interventions (e.g., 1 session delivered in the hospital, first contact was beyond 1 week, school-based interventions, psychotherapy, pharmacotherapy), and noncomparative study designs (e.g., single group design, case report). In terms of the comprehensiveness of the literature search, most SRs did not have a restriction on publication date, with 1 including studies from January 1, 2000, to the date of the search. The end search dates ranged from January 2015 to May 31, 2021. The 2 SRs with relevant primary studies (Witt et al. and Chaudhary et al.) each included 1 primary study relevant to this review. These 2 SRs will be described further in subsequent sections of this review, with the exception of the Summary of Critical Appraisal, which will discuss all 6 included SRs.

Additional details regarding the characteristics of included publications are provided in Table 2.

Study Design
The SRs were published in 2021 and 2020. Date ranges covered by their searches were up to July 4, 2020 and January 2019. The SR by Witt et al. included RCTs and the SR by Chaudhary et al. included RCTs and nonrandomized studies. There was no overlap in the relevant studies for these reviews because the SR by Witt et al. included an RCT and the SR by Chaudhary et al. included a nonrandomized clinical trial.

Country of Origin
The first authors for the 2 SRs were from Australia and Pakistan. Both primary studies were conducted in the US.

Patient Population
In the relevant primary studies in both SRs, the populations were adolescents who attempted suicide and were evaluated in the emergency department. The primary study in the SR by Witt et al. included participants who received medical attention in emergency departments and pediatric wards, although it is unclear what proportion were cared for in wards.
Interventions and Comparators
In the SR by Witt et al.,\textsuperscript{15} the relevant RCT evaluated an intervention described as a “compliance-enhanced intervention,” which included a single 1-hour session, delivered in the emergency department before discharge, to review expectations for outpatient treatment, address common treatment misconceptions, a verbal treatment contract, and a series of 4 telephone calls at 1, 2, 4, and 8 weeks after discharge (duration not reported). The intervention was delivered by 3 postdoctoral fellows in psychology (experience not reported). Treatment as usual (standard disposition planning) consisted of a “brief” (duration not reported) inpatient treatment and/or outpatient appointment, as appropriate.

In the SR by Chaudhary et al.,\textsuperscript{16} the relevant nonrandomized clinical trial evaluated an intervention that included 3 phone interviews over 8 weeks that were scheduled with patients and parents for 1, 2, and 6 weeks after discharge. These phone calls focused on treatment expectations, outpatient services, problems, concerns, and resistance of patients and their parents to attend predecided outpatient psychotherapy sessions. The intervention was delivered by a doctor-level clinician. The comparison group was described as treatment as usual, without further details.

Outcomes
Outcomes reported in the relevant primary studies within the SRs include:

- psychotherapy session “no shows”\textsuperscript{16}
- completed the full course of treatment\textsuperscript{15}
- average number of sessions attended\textsuperscript{15,16}
- suicide deaths\textsuperscript{15}
- suicide reattempts\textsuperscript{16}
- repetition of self-harm at 6 months follow-up\textsuperscript{15}

In the SR by Witt et al.,\textsuperscript{15} repetition of self-harm was identified through self-report, collateral report, clinical records, or research monitoring systems, and suicide was identified through register-recorded deaths or reports from collateral informants, such as family members or neighbours. Depression, hopelessness, general functioning, social functioning, and suicidal ideation were also reported in the SR by Witt et al.,\textsuperscript{15} but with no data available.

Chaudhary et al.\textsuperscript{16} did not report on the outcomes of interest in the inclusion criteria, but outcomes related to suicide and self-harm were captured in the results tables (e.g., suicidal ideations, repetition of suicidal behaviour, rate of suicidal attempts after discharge from the hospital, completion of suicide, repetition of deliberate self-harm within 6 months of the initial incident, number of self-harm episodes after receiving treatment, readmission for self-poisoning), depression, hopelessness, social and psychological well-being, and outpatient services outcomes (e.g., engagement with outpatient services).
Summary of Critical Appraisal

Systematic Reviews
The SRs were assessed using AMSTAR 2. Among the 2 SRs with relevant primary studies, the SR by Witt et al. is a Cochrane review and was well conducted and reported, with no major limitations identified. Chaudhary et al. searched 2 electronic databases, did not restrict inclusion of studies based on language, provided a PRISMA flow diagram, and performed study selection and data extraction with 2 reviewers; however, there were several limitations in conduct and reporting. There was no mention of a registered protocol, it was unclear if supplemental searching was performed, elements of PICO were not well described in the eligibility criteria or for the included studies, a list of excluded studies was not provided, risk of bias was not formally done with a tool, there was no mention of assessment for publication bias, and the source of funding of the included primary studies and for the review itself was not reported.

There were strengths across the 4 SRs with no relevant primary studies. All searched 2 or more databases, provided a PRISMA flow diagram, reported declarations and conflicts of interest, and described the statistical analyses. However, there were differences in limitations in quality of conduct and reporting across the 4 SRs. Some quality criteria were not conducted or reported: publications were restricted to English publications, elements of PICO for the inclusion criteria were not well described, no details on the conduct for study selection and data extraction were provided, a list of excluded studies was not provided (with varying levels of details for exclusion in the PRISMA flow diagram), no details around how risk of bias was performed, some PICO elements of the included studies were not well described, it was unclear if it was assessed, no details of funding for the included studies in the SR, and no details of funding source for the SR. It is possible that some studies included in the SRs met our inclusion criteria, but we were unable to determine this due to lack of details.

Additional details regarding the strengths and limitations of included publications are provided in Table 3.

Summary of Findings

Clinical Effectiveness of Active Contact and Follow-Up Interventions
Six SRs were identified that met the inclusion criteria for this research question, of which 2 contained relevant primary studies: 1 RCT from the SR by Witt et al. and 1 nonrandomized clinical trial from the SR by Chaudhary et al. There was no overlap of the relevant primary studies included in the SRs.

Adherence to Psychotherapy
Overall, there may be no difference in adherence to psychotherapy between those who received telephone follow-up care and those who received treatment as usual.

Evidence from 1 RCT in the SR by Witt et al. reported no significant difference in those patients who completed the full course of treatment for of the compliance-enhancement intervention compared with those who received treatment as usual (odds ratio [OR] = 1.59; 95% CI, 0.59 to 4.33). Evidence from 1
nonrandomized clinical trial identified in the SR by Chaudhary et al.\textsuperscript{16} reported psychotherapy “no shows” of 9% for those receiving telephone calls and 18% for those receiving treatment as usual (no statistical comparison reported).

Both primary studies in the SRs\textsuperscript{15,16} reported on the average number of sessions. The RCT in the SR by Witt et al.\textsuperscript{15} reported no difference in the average number of sessions attended between groups (mean difference = 1.30; 95% CI, −1.28 to 3.88), whereas the nonrandomized trial in the SR by Chaudhary et al.\textsuperscript{16} reported the mean number of sessions was 5.5 in the group receiving telephone calls compared with 3.9 in the treatment as usual group (no statistical comparison reported).

**Number of Reattempts and Death by Suicide**

Overall, there may be no difference in the number of reattempts or death by suicide between those who received telephone follow-up care and those who received treatment as usual.

The RCT in the SR by Witt et al.\textsuperscript{15} reported no deaths by suicide in either group by the final follow-up assessment. Additionally there was no significant difference between groups in patients repeating self-harm at 6 month follow-up (OR = 0.67; 95% CI, 0.15 to 3.08).\textsuperscript{15} Evidence from the nonrandomized trial in the SR by Chaudhary et al.\textsuperscript{16} reported that the group receiving telephone calls had 0% reattempted suicide, whereas the treatment as usual group had a 9% suicide reattempt rate.

**Other Outcomes**

Witt et al.\textsuperscript{15} sought evidence regarding depression, hopelessness, general functioning, social functioning, and suicidal ideation; no evidence was identified for these outcomes in the primary study relevant to this review. Table 4 presents the main study findings.

**Clinical Effectiveness of Time Durations for Follow-Up Care**

No relevant evidence regarding the time durations for initiation of follow-up care for children and adolescents who presented to emergency care for suicide attempts or suicide ideations was identified; therefore, no summary can be provided.

**Evidence-Based Guidelines About Timing, Modality, and Health Care Professional Involved in Follow-Up Care**

No relevant evidence-based guidelines regarding the timing, modality, and health care professionals involved in follow-up care for children and adolescents who present to emergency care for suicide attempts or suicide ideations was identified; therefore, no summary can be provided.

**Limitations**

There were some limitations to the SRs that would prevent a definitive conclusion on clinical effectiveness of active contact and follow-up intervention for children and adolescents who present to emergency care for suicide attempts or suicide ideations.
Six SRs were included with inclusion criteria broader than the scope of this review, particularly around the populations and interventions included in the primary studies. In most reviews, there was poor reporting of the patient population and interventions, making it difficult to determine if primary studies were relevant to this review. Therefore, it is possible that some primary studies within the SRs were missed in this review (although a search of the abstract for the primary studies did not result in any studies that would meet the eligibility criteria of this review). Two SRs provided sufficient details to identify 2 relevant primary studies. The SR by Witt et al.\textsuperscript{15} was well conducted and reported; however, the relevant primary study was judged on 6 domains of risk of bias, with 3 judged as having “some concerns” and 3 judged as “high risk.” The quality of conduct and reporting for the SR by Chaudhary et al.\textsuperscript{16} was poor, and they did not perform a formal risk of bias assessment, with the limitation for the relevant primary study listed as “small sample size.”

Both relevant primary studies in the SRs were conducted in the US (1 in Northeast US, and the other did not provide any further details), with data from 1 emergency department (no further details about the setting reported), and sample sizes of 64 participants\textsuperscript{15} and 97 participants\textsuperscript{16}. These limit the generalizability because the capacity of emergency departments may differ across regions and countries and the participants in these studies may not be representative of all children and adolescents who present to emergency departments for suicidal ideation or attempts. The intervention in both primary studies evaluated telephone follow-up, which limits the generalizability of follow-up aftercare for other modalities (e.g., email, text message, home visit). Neither primary study in the SRs reported on mental health outcomes (e.g., depression, anxiety) or harms of the intervention.

No studies were identified that evaluated different time durations from emergency department discharge for follow-up care. Although the 2 relevant primary studies in the SRs reported on a different numbers of telephone calls, with the first call within 1 week of discharge, they were compared with treatment as usual, which was not reported or was missing details (i.e., duration), not with a different time duration.

No evidence-based guidelines that met our inclusion criteria were identified. One potential guideline by the National Institute for Health and Care Excellence (NICE) provided a recommendation on first point of contact within 48 hours after discharge,\textsuperscript{7} but was excluded because the recommendation was not specific to the child and adolescent population. This guideline has been listed in Appendix 5, Table 5.

**Conclusions and Implications for Decision- or Policy-Making**

The evidence is scarce about active contact and follow-up care in children and adolescents (younger than 18 years) who present to the emergency department with attempted suicide or suicide ideation but are not subsequently admitted. This review included 6 SRs,\textsuperscript{8,9,11,12,15,16} and the included primary studies were published from 1976 to 2021. Two SRs\textsuperscript{15,16} each identified 1 relevant primary study that evaluated telephone follow-up care; these primary studies were published in 1997 and 2002. Although other active follow-up care interventions were identified in the SRs, poor reporting of the PICO elements of the included studies limited the ability to identify other active follow-up interventions (e.g., email, text message, home visit) or relevant studies conducted in the past 20 years. When PICO elements were not clearly described within the SRs, an
additional search for the abstracts was performed, which did not reveal any further relevant studies, although abstracts do not always sufficiently describe the PICO elements. There were no studies evaluating different time durations for follow-up care. An SR of primary studies using the same inclusion criteria as this Rapid Review may identify additional relevant studies because the population and intervention within the primary study would likely provide sufficient details to determine eligibility.

Although the 2 relevant primary studies within the SRs both evaluated telephone follow-up, there was a difference between the interventions. The compliance-enhancement approach included an additional 1-hour session before emergency department discharge and 1 additional telephone call. Evidence from the RCT evaluating the compliance-enhancement approach, included in the SR by Witt et al., showed no significant differences between participants who received the single-session and follow-up telephone calls and participants who received treatment as usual in the number of deaths by suicide, those who completed the full course of treatment, or the average number of sessions attended. Chaudhary et al. reported means and rates but did not state whether there was a statistical difference between groups. The limitations of the included literature should be considered when interpreting the findings of this report. Although the SR by Witt et al. was well conducted and reported, the relevant primary study was at high risk of bias. Additionally, both relevant primary studies in the SRs had small sample sizes (N = 64 and N = 97), which affects the number of possible events, and the lack of observed significant differences could be due to sample size rather than lack of intervention effect.

Several reviews and primary studies have evaluated different modalities of follow-up care in adults, participants admitted to hospitals, or opportunistic screening of adolescents presenting to the emergency department for nonpsychiatric reasons, or with no comparison group. Although these sources generally indicated that active follow-up is associated with positive outcomes, such as reducing the risk of repeat suicide attempts and increasing connectedness to ongoing care for adults, it is unclear if these populations are generalizable to children and adolescents presenting to the emergency department due to suicidal ideations or attempts.

We did not identify any evidence-based guidelines regarding the timing, modality, or health care professionals involved in the follow-up care for people younger than 18 years who present to emergency care for suicide attempts or suicide ideations. However, guidance documents and guidelines for adults (refer to Appendix 5, Table 5) generally indicate that follow-up should occur within 48 hours, particularly for adults with safety concerns of subsequent self-harm. These guidance documents also suggest different modalities for follow-up during that time, including telephone calls, visits, and electronic communication. Given the higher risk and potential vulnerability experienced by children and adolescents, in the absence of formal clinical guidelines, ensuring person-centred follow-up care for all children and adolescents is provided within 48 hours may be considered, similar to adults presenting with high concerns. As the number of children and adolescents presenting to emergency departments for suicide attempts has increased since the COVID-19 pandemic, future research in this area should evaluate all modalities for active follow-up care, including text messages, telephone calls, email, and home visits specific to children and adolescents presenting to the emergency department due to suicidal ideation or attempt who are not admitted to hospital.
References


Appendix 1: Selection of Included Studies

Figure 1: Selection of Included Studies

515 citations identified from electronic literature search and screened

489 citations excluded

26 potentially relevant articles retrieved for scrutiny (full text, if available)

14 potentially relevant reports retrieved from other sources (grey literature, handsearch)

40 potentially relevant reports

34 reports excluded:
- irrelevant population (12)
- irrelevant intervention (13)
- other (review articles, editorials) (9)

6 reports included in review
Appendix 2: Characteristics of Included Publications

Table 2: Characteristics of Included Systematic Reviews

<table>
<thead>
<tr>
<th>Study citation, country, funding source</th>
<th>Study designs and numbers of primary studies included</th>
<th>Population characteristics</th>
<th>Intervention and comparator(s)</th>
<th>Clinical outcomes, length of follow-up</th>
</tr>
</thead>
<tbody>
<tr>
<td>Skopp et al. (2023)¹¹</td>
<td>RCTs Studies published up to February 11, 2020</td>
<td>Broader review: Patients deemed by medical or behavioural health services to be at elevated risk for suicide. All included studies were in adults, with some studies including patients discharged from psychiatric care.</td>
<td>Eligible intervention: Caring Contacts (contacts were text-based, including email, letter, text message, or postcard). Most studies had follow-up care starting more than 1-week post emergency department discharge. Comparator: Not specified</td>
<td>Outcomes reported: All-cause mortality; suicide mortality; suicidal ideation; suicide attempts; emergency department readmissions; psychiatric inpatient and general admissions; representation for self-harm; medial evacuation; self-injury Follow-up: NR</td>
</tr>
<tr>
<td>Hou et al. (2022)⁰¹</td>
<td>Classification of social support interventions portion of review: all studies Meta-analysis: RCTs Studies published up to May 31, 2021 Included studies: 77 studies in classification review; 14 RCTs included in meta-analysis; 0 relevant to the present review</td>
<td>Broader review: Suicidal individuals Included studies were in adults, patients in psychiatric inpatient units, in hospital, or delivered to students in a school setting.</td>
<td>Eligible intervention: Studies aimed at preventing suicide through method(s) that directly provide social support. Many studies had follow-up care starting more than 1-week post emergency department discharge. Comparator: Treatment as usual or “the latter serving group” for comparative studies; No comparator for noncomparative studies</td>
<td>Outcomes reported: Suicide; suicide attempt; social support-related outcome Follow-up: NR</td>
</tr>
<tr>
<td>Witt et al. (2021)¹⁵</td>
<td>RCTs Studies published up to July 4, 2020</td>
<td>Broader review: Children and adolescents (up to 18 years of age) with a recent (within 6 months of trial entry) presentation to hospital or clinical service for self-harm</td>
<td>Eligible interventions: Psychosocial or pharmacological treatments. Many studies evaluated cognitive-behavioural therapy, group therapy, dialectical behaviour therapy, mentalization based therapy, home-</td>
<td>Outcomes reported in relevant primary study: adherence to psychotherapy; repeat self-harm and death by suicide Follow-up: Up to 2 years</td>
</tr>
<tr>
<td>Study citation, country, funding source</td>
<td>Study designs and numbers of primary studies included</td>
<td>Population characteristics</td>
<td>Intervention and comparator(s)</td>
<td>Clinical outcomes, length of follow-up</td>
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<tr>
<td>Chaudhary et al. (2020)(^6) Pakistan</td>
<td>RCTs, nonrandomized clinical trials, cohort, descriptive analysis Studies published up to January 2019</td>
<td>(e.g., self-poisoning, self-injury) Some included studies had patients admitted to hospital. <strong>Relevant included study:</strong> Adolescents aged 12 to 18 years old presenting to hospital following a suicide attempt</td>
<td>based family therapy, and pharmacological therapies. <strong>Comparator:</strong> Treatment as-usual, routine psychiatric care, enhanced usual care, active comparator, placebo, alternative pharmacological treatment, or a combination of these. <strong>Relevant intervention:</strong> Described as compliance-enhancement approach; a single 1-hour session, before emergency department discharge, to review expectations for outpatient treatment, address common treatment misconceptions, a verbal treatment contract, and a series of 4 telephone calls at 1, 2, 4 and 8 weeks' post-discharge (duration not reported). Intervention delivered by 3 postdoctoral fellows in psychology (experience not reported) <strong>Relevant comparator:</strong> Treatment as usual, a standard disposition planning consisting of a brief (duration not reported) inpatient treatment and/or an outpatient appointment, as appropriate</td>
<td><strong>Outcomes reported in relevant primary study:</strong> School functioning; Suicidal behaviours, Psychotherapy visits <strong>Follow-up:</strong> NR</td>
</tr>
<tr>
<td>for Health Research</td>
<td>RCTs; 1 RCT relevant to the present review</td>
<td>Broader review: No restriction on race, place, sex, age, and ethnicity were applied. Included studies were in adults, patients admitted to hospital, or patients in psychiatric inpatient units <strong>Relevant included study:</strong> Adolescents aged 12 to 18 years old seen at general hospital for a</td>
<td>Intervention: Interventions to target suicidal behaviours after discharge from a medical facility to the community Included studies had follow-up care starting more than 1-week post emergency department discharge, and sent letters to general practitioners, <strong>Comparator:</strong> NR <strong>Relevant intervention:</strong> Structured 3 telephone intervention with patients and parents at 1, 2,</td>
<td></td>
</tr>
<tr>
<td>Study citation, country, funding source</td>
<td>Study designs and numbers of primary studies included</td>
<td>Population characteristics</td>
<td>Intervention and comparator(s)</td>
<td>Clinical outcomes, length of follow-up</td>
</tr>
<tr>
<td>----------------------------------------</td>
<td>------------------------------------------------------</td>
<td>-----------------------------</td>
<td>---------------------------------</td>
<td>----------------------------------------</td>
</tr>
</tbody>
</table>
| Doupnik et al. (2020)  
US  
Funding source: National Institute of Mental Health of the National Institutes of Health | Studies with a comparison group  
Studies published January 1, 2000 to December 31, 2019  
Included studies: 14 studies (design not reported); 0 relevant to the present review | Suicide attempt requiring medical care | and 6 weeks following discharge; Intervention delivered by a doctor-level clinician  
**Relevant comparator:** Treatment as usual (not further described) | Outcomes listed in PROSPERO: Suicide death; Suicide ideation; Suicide attempt; Suicide-related health care visit  
Outcomes included in meta-analysis: Subsequent suicide attempts; Linkage to follow-up care; Depression symptoms at follow-up  
Follow-up: NR |
| Inagaki et al. (2019)  
Japan  
Funding source: Japan Agency for Medical Research and Development | RCTs  
Studies published up to January 2015  
Included studies: 28 RCTs (14 RCTs for active follow-up); 0 relevant to the present review | Participants had attempted suicidal behaviour within 1 month and had been admitted to an emergency department for their suicidal behaviour  
Most included studies were in adults. | All interventions (e.g., psychosocial intervention, psychotherapy, pharmacotherapy).  
Included studies had follow-up care starting more than 1-week post emergency department discharge, continuous case management, and pharmacotherapy  
**Comparator:** Placebo, treatment as usual and other expected intervention  
**Relevant interventions:** Interventions in the trial was performed while the patients were in the emergency department or a subsequent ward; Active contact and follow-up interventions (intensive care plus outreach, brief interventions and contact, letter/postcard, telephone, and composite of letter/postcard and telephone) | Outcomes reported: Recurrence of attempted suicide or self-harm; Completed suicide; Any cause of death  
Follow-up: NR |

NR = not reported; RCT = randomized controlled trial.
Note this appendix has not been copy-edited.
### Table 3: Strengths and Limitations of Systematic Reviews Using AMSTAR 2\textsuperscript{13}

<table>
<thead>
<tr>
<th>Strengths</th>
<th>Limitations</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Skopp et al. (2023)\textsuperscript{11}</strong></td>
<td><strong>Limitations</strong></td>
</tr>
<tr>
<td>• A protocol was registered on the PROSPERO website</td>
<td>• Some elements of PICO were not well described in the eligibility criteria (e.g., population, comparators, outcome)</td>
</tr>
<tr>
<td>• Four electronic databases were searched; supplemental searching performed</td>
<td>• Restriction on language of inclusion (English only)</td>
</tr>
<tr>
<td>• Study selection performed independently by 2 reviewers, with disagreements adjudicated by consensus agreement</td>
<td>• A list of excluded studies was not provided, but high-level reasons were provided in the PRISMA flow diagram</td>
</tr>
<tr>
<td>• A PRISMA flow diagram was provided (supplemental materials)</td>
<td>• Risk of bias assessed, but no details on how this was performed</td>
</tr>
<tr>
<td>• Data extraction performed by 2 reviewers</td>
<td>• Certainty of the evidence for some outcomes was provided, but unclear if publication bias assessed within</td>
</tr>
<tr>
<td>• Statistical analysis well described</td>
<td>• Source of funding of the included primary studies not provided</td>
</tr>
<tr>
<td>• PICO elements of primary studies sufficiently described</td>
<td>• No details on funding for the review was reported</td>
</tr>
<tr>
<td>• The authors state they have no conflict of interest to disclose</td>
<td></td>
</tr>
</tbody>
</table>

| **Hou et al. (2022)\textsuperscript{12}** | |
| • A protocol was registered on the PROSPERO website | • Restriction on language of inclusion (English only) |
| • Six electronic databases were searched; supplemental searching performed | • Some elements of PICO were not well described in the eligibility criteria (e.g., population) |
| • Study selection performed independently by 5 reviewers (title/abstract) or 2 reviewers (full text), with disagreements resolved via face to face meetings | • A list of excluded studies was not provided, but high-level reasons were provided in the PRISMA flow diagram |
| • A PRISMA flow diagram was provided | • Some elements of PICO were not well described for the included studies (e.g., population, comparator) |
| • Data extraction independently performed by 2 reviewers extracted data, with disagreements resolved by consensus | • Source of funding of the included primary studies not provided |
| • Risk of bias assessed independently by 2 reviewers with disagreement resolved by consensus | |
| • Statistical analysis well described | |
| • Publication bias assessed | |
| • Authors declared no competing interests | |

<p>| <strong>Witt et al. (2021)\textsuperscript{15}</strong> | |
| • No formal statement around a protocol, however Cochrane Reviews begin with a protocol, and there is a statement that there were no differences between the protocol and the full review | • No major limitations identified. |
| • Six electronic databases were searched; supplemental searching performed | |
| • Elements of PICO were well described for inclusion | |
| • No restrictions on language of inclusion | |
| • Study selection performed independently by 2 reviewers, with | |</p>
<table>
<thead>
<tr>
<th>Strengths</th>
<th>Limitations</th>
</tr>
</thead>
<tbody>
<tr>
<td>disagreements resolved in consultation with the senior review author</td>
<td>No mention of a protocol</td>
</tr>
<tr>
<td>• A PRISMA flow diagram was provided</td>
<td>• Unclear if supplemental searching was performed</td>
</tr>
<tr>
<td>• Two reviewers extracted data, with disagreements resolved by consensus</td>
<td>• Some elements of PICO were not well described in the eligibility criteria (e.g., vague details around relevant interventions, no information on relevant comparators)</td>
</tr>
<tr>
<td>• Risk of bias assessed independently by 2 reviewers with disagreement resolved by consensus</td>
<td>• Some elements of PICO of the included studies were not well described (e.g., population)</td>
</tr>
<tr>
<td>• Statistical analysis well described</td>
<td>• A list of excluded studies was not provided, but high-level reasons were provided in the PRISMA flow diagram</td>
</tr>
<tr>
<td>• PICO elements of primary studies well described</td>
<td>• Risk of bias not formally done with a tool, with minimal reporting of potential bias under limitations for each included study</td>
</tr>
<tr>
<td>• A list of excluded studies and reason for exclusion was provided</td>
<td>• No mention of assessment for publication bias</td>
</tr>
<tr>
<td>• Certainty of the evidence for some outcomes was provided, with publication bias assessed within</td>
<td>• Source of funding of the included primary studies not provided</td>
</tr>
<tr>
<td>• Declarations of interest were reported</td>
<td>• No details on funding for the review was reported</td>
</tr>
<tr>
<td>• Sources of funding for the review and for the included studies within the review was provided</td>
<td></td>
</tr>
</tbody>
</table>

**Chaudhary et al. (2020)\(^6\)**

- Two electronic databases were searched
- No restrictions on language of inclusion
- A PRISMA flow diagram was provided
- Study selection performed independently by 2 reviewers, with conflicts resolved by consensus or with a third author
- Two reviewers extracted data, with data cross-checked for accuracy by senior author
- Authors report that there were no conflicts of interest

**Doupnik et al. (2020)\(^8\)**

- A protocol was registered on the PROSPERO website
- Five electronic databases were searched; supplemental searching performed
- Study selection performed independently by 2 reviewers, with team meetings to discuss and resolve discrepancies and reach consensus on all inclusion decisions
- A PRISMA flow diagram was provided
- Data extraction performed independently by 2 reviewers (no details on how disagreements were resolved)
- Statistical analysis well described

- Restriction on language of inclusion (English only)
- A list of excluded studies was not provided, no additional details in PRISMA flow diagram
- Risk of bias assessments performed, but no details on the conduct
- Some elements of PICO of the included studies were not well described (e.g., comparator, study design)
- Source of funding of the included primary studies not provided
<table>
<thead>
<tr>
<th>Strengths</th>
<th>Limitations</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Small study effects (e.g., publication bias) assessed</td>
<td>• Restriction on language of inclusion (English only)</td>
</tr>
<tr>
<td>• Sources of funding for the review was provided</td>
<td>• No details on the conduct for study selection and data extraction</td>
</tr>
<tr>
<td>• Authors provided conflict of interest statements</td>
<td>• Risk of bias assessments performed, but no details on the conduct</td>
</tr>
<tr>
<td></td>
<td>• A list of excluded studies was not provided, no additional details in PRISMA flow diagram</td>
</tr>
<tr>
<td></td>
<td>• Some elements of PICO of the included studies were not well described (e.g., population)</td>
</tr>
<tr>
<td></td>
<td>• Source of funding of the included primary studies not provided</td>
</tr>
</tbody>
</table>

Inagaki et al. (2019)²

- A protocol was registered on the PROSPERO website
- Four electronic databases were searched; supplemental searching performed
- Elements of PICO were well described for inclusion
- A PRISMA flow diagram was provided
- Statistical analysis well described
- Publication bias assessed
- Authors declared conflicts of interest
- Sources of funding for the was provided

AMSTAR 2 = A Measurement Tool to Assess Systematic Reviews 2; PICO = population, intervention, comparator, outcomes; PRISMA = Preferred Reporting Items for Systematic Reviews and Meta-Analyses.
## Appendix 4: Main Study Findings

Note this appendix has not been copy-edited.

### Table 4: Summary of Findings by Outcome

<table>
<thead>
<tr>
<th>Variable</th>
<th>Witt et al. (2021)(^{15})</th>
<th>Chaudhary et al. (2020)(^{16})</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N = 64</td>
<td>N = 97</td>
</tr>
<tr>
<td><strong>Adherence to psychotherapy</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Psychotherapy session “no shows”</td>
<td>NR</td>
<td>Intervention: 9%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Comparison: 18%</td>
</tr>
<tr>
<td>Completed the full course of treatment</td>
<td>Intervention: 17/29</td>
<td>NR</td>
</tr>
<tr>
<td></td>
<td>Comparison: 16/34</td>
<td></td>
</tr>
<tr>
<td></td>
<td>OR 1.59, 95% CI 0.59 to 4.33</td>
<td></td>
</tr>
<tr>
<td>Average number of sessions attended</td>
<td>Intervention: mean 7.70 sessions (SD 5.80, n = 29)</td>
<td>Intervention: mean 5.5 sessions (SD 4.40, n = 34)</td>
</tr>
<tr>
<td></td>
<td>Comparison: mean 6.40 sessions (SD 4.40, n = 34)</td>
<td>Comparison: mean 3.9 session</td>
</tr>
<tr>
<td></td>
<td>MD 1.30 95% CI −1.28 to 3.88; N = 63</td>
<td></td>
</tr>
<tr>
<td><strong>Number of reattempts and death by suicide</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Suicide deaths</td>
<td>Intervention: 0 suicides</td>
<td>NR</td>
</tr>
<tr>
<td></td>
<td>Comparison: 0 suicides</td>
<td></td>
</tr>
<tr>
<td>Suicide reattempts</td>
<td>NR</td>
<td>Intervention: 0% reattempted</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Comparison: 9% reattempted</td>
</tr>
<tr>
<td>Repetition of self-harm at 6 months follow-up</td>
<td>Intervention: 3/29</td>
<td>NR</td>
</tr>
<tr>
<td></td>
<td>Comparison: 5/34</td>
<td></td>
</tr>
<tr>
<td></td>
<td>OR 0.67, 95% CI 0.15 to 3.08</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Repeat self-harm episodes:</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Intervention vs comparison:</td>
<td></td>
</tr>
<tr>
<td></td>
<td>mean 0.10 vs. 0.15</td>
<td></td>
</tr>
<tr>
<td><strong>Depression</strong></td>
<td>NA</td>
<td>No data available</td>
</tr>
<tr>
<td></td>
<td></td>
<td>NR</td>
</tr>
<tr>
<td><strong>Hopelessness</strong></td>
<td>NA</td>
<td>No data available</td>
</tr>
<tr>
<td></td>
<td></td>
<td>NR</td>
</tr>
<tr>
<td><strong>General functioning</strong></td>
<td>NA</td>
<td>No data available</td>
</tr>
<tr>
<td></td>
<td></td>
<td>NR</td>
</tr>
<tr>
<td><strong>Social functioning</strong></td>
<td>NA</td>
<td>No data available</td>
</tr>
<tr>
<td></td>
<td></td>
<td>NR</td>
</tr>
<tr>
<td><strong>Suicidal ideation</strong></td>
<td>NA</td>
<td>No data available</td>
</tr>
</tbody>
</table>

CI = confidence interval; MD = mean difference; N = number; NA = not applicable; NR = not reported; OR = odds ratio; SD = standard deviation.
Appendix 5: References and Guidance of Potential Interest

Note this appendix has not been copy-edited.

The subsequent list of potentially relevant articles includes studies that may be relevant but were not included in the main analysis as they did not fulfill the PICO criteria (primarily due to age of study population). However, note that interventions assessing follow-up care among adults was not within the scope of the literature search and therefore, there are likely other studies related to adult population not mentioned here.

Korczak et al. (2020) details an ongoing RCT in Canada about a youth specific suicide prevention intervention. Although the primary aim of the study is to assess the specific intervention within emergency care, monitoring the study results may be useful for developing follow-up protocols.

Informal Scan of Emergency Department Follow-Up Guidance or Protocols for Suicide Ideation or Attempts

Online scanning and grey literature searching identified the following publicly available guidance documents or resources related to follow-up care produced by jurisdictions in Canada or internationally. The scan was not systematic or comprehensive. In particular, Canadian jurisdictions and health care centres may have other resources that are not publicly or easily accessible. The scan was also limited to English-language resources.

Table 5 lists the identified guidance documents, protocols, or recommendations that mention follow-up care among other aspects of delivering care for self-harm or suicide prevention. The recommendations summarized in the table are specifically related to follow-up care for people who are not admitted into inpatient settings and have been discharged from emergency care or other acute settings. These resources generally indicate that active follow-up care is recognized as being an important part of the care pathway. The resources recommend different options for follow-up periods, modalities, and health care professionals, but the resources do not consistently provide specific details about modalities or health care professionals who should be involved. Some reports recommend multiple stages of follow-up (e.g., immediate follow-up and subsequent follow-up over a longer time period). Guidance specifically for children and adolescents (under 18 years) was not identified.

Table 5: Guidance Documents Describing Follow-Up Protocols for People Visiting Emergency Care for Suicide Attempts or Ideation

<table>
<thead>
<tr>
<th>Jurisdiction</th>
<th>Organization or hospital</th>
<th>Recommendations or findings</th>
<th>Link</th>
</tr>
</thead>
</table>
| Canada (pan-Canadian) | Mental Health Commission of Canada | • Follow-up plans for people attending emergency care should include contacting the patient 24 to 72 hours after discharge.  
• Contact can be made by phone, text message, letter or postcard, home visit, and/or email) and be initiated by a volunteer, a nonspecialized | Post-Attempt Followup and Regular Contact Interventions Evidence Brief on Suicide Care (publication date not clear) |
<table>
<thead>
<tr>
<th>Jurisdiction</th>
<th>Organization or hospital</th>
<th>Recommendations or findings</th>
<th>Link</th>
</tr>
</thead>
</table>
| Canada (British Columbia) | BC Mental Health and Addiction Services | • Any health care provider serving an individual who is suicidal to follow up within 24 hours (or sooner if required).  
• In-person follow-up within 7 days (or sooner if required) of discharge from inpatient services for those with recent suicidal behaviour.  
• Patients should be assessed and carefully monitored within the first 30 days following discharge from inpatient services. | The Provincial Suicide Clinical Framework (January 2011) |
| UK | The National Institute for Health and Care Excellence | Regarding initial after care following self-harm, guidelines recommend:  
• The mental health team, general practitioner, or care providers should provide initial aftercare within 48 hours of presentation.  
• This immediate aftercare is intended for people who may have ongoing safety concerns.  
• The committee which developed the guidelines, stated that the 48 hours aftercare would likely be beneficial for everyone, but may not be feasible in all settings, hence why the criteria for ‘ongoing safety concerns’ was included. | Self-harm: assessment, management and preventing recurrence (September 2022) |
| Australia | Black Dog Institute | A member of the suicide response team within emergency or acute care should ensure people:  
• Receive a follow-up referral and appointment within 24 hours following discharge.  
• Receive a DCP  
• The DCP should be provided to the person in written format, verbally, and in any other format preferred by the person (email, mail).  
• The health care team should also provide a copy of the DCP to the person's general practitioner or aftercare service within 24 hours of the discharge, and, with the person's consent, to their carer, family, or friends. | Guidelines for integrated suicide-related crisis and follow-up care in Emergency Departments and other acute settings (November 2017) |
| US (California) | Mental Health Services Oversight and Accountability Commission | California has set an objective for all hospitals and emergency departments to:  
• Develop protocols and policies to have a goal of connecting individuals being discharged from the emergency departments or inpatient services to outpatient services within 24 to 48 hours.  
• Follow-up care must be linguistically and culturally respectful. | California’s Strategic Plan for Suicide Prevention 2020 – 2025 (December 2019) |
### Jurisdiction

<table>
<thead>
<tr>
<th>Jurisdiction</th>
<th>Organization or hospital</th>
<th>Recommendations or findings</th>
<th>Link</th>
</tr>
</thead>
<tbody>
<tr>
<td>US (New York State)</td>
<td>NY Center for Practice Innovations</td>
<td>• 2 or more follow-ups, 1 of which should be within discharge of 24 to 72 hours after discharge.</td>
<td>A Guide for Clinicians (2021)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• 2 or more follow-ups are recommended, 1 within 24 to 72 hours of discharge, and 1 following the first outpatient appointment.</td>
<td></td>
</tr>
</tbody>
</table>

DCP = discharge care plan.

### Potentially Relevant Articles With Adult or Unclear Population


### Additional References

