JAMA | US Preventive Services Task Force | EVIDENCE REPORT

Screening for Depression and Suicide Risk in Children and Adolescents Updated Evidence Report and Systematic Review for the US Preventive Services Task Force

Meera Viswanathan, PhD; Ina F. Wallace, PhD; Jennifer Cook Middleton, PhD; Sara M. Kennedy, MPH; Joni McKeeman, PhD; Kesha Hudson, PhD; Caroline Rains, MPH; Emily B. Vander Schaaf, MD, MPH; Leila Kahwati, MD, MPH

IMPORTANCE Depression, suicidal ideation, and self-harm behaviors in youth are associated with functional impairment and suicide.

OBJECTIVE To review the evidence on screening for depression or suicide risk in children and adolescents to inform the US Preventive Services Task Force (USPSTF).

DATA SOURCES PubMed, Cochrane Library, PsycINFO, CINAHL, and trial registries through July 19, 2021; references, experts, and surveillance through June 1, 2022.

STUDY SELECTION English-language, randomized clinical trials (RCTs) of screening for depression or suicide risk; diagnostic test accuracy studies; RCTs of psychotherapy and first-line pharmacotherapy; RCTs, observational studies, and systematic reviews reporting harms.

DATA EXTRACTION AND SYNTHESIS Two reviewers assessed titles/abstracts, full-text articles, and study quality and extracted data; when at least 3 similar studies were available, meta-analyses were conducted.

MAIN OUTCOMES AND MEASURES Test accuracy, symptoms, response, remission, loss of diagnosis, mortality, functioning, suicide-related events, and adverse events.

RESULTS Twenty-one studies (N = 5433) were included for depression and 19 studies (N = 6290) for suicide risk. For depression, no studies reported on the direct effects of screening on health outcomes, and 7 studies (n = 3281) reported sensitivity of screening instruments ranging from 0.59 to 0.94 and specificity from 0.38 to 0.96. Depression treatment with psychotherapy was associated with improved symptoms (Beck Depression Inventory pooled standardized mean difference, -0.58 [95% CI, -0.83 to -0.34]; n = 471; 4 studies; and Hamilton Depression Scale pooled mean difference, -2.25 [95% CI, -4.09 to -0.41]; n = 262; 3 studies) clinical response (3 studies with statistically significant results using varying thresholds), and loss of diagnosis (relative risk, 1.73 [95% CI, 1.00 to 3.00; n = 395; 4 studies). Pharmacotherapy was associated with improvement on symptoms (Children's Depression Rating Scale-Revised mean difference, -3.76 [95% CI, -5.95 to -1.57; n = 793; 3 studies), remission (relative risk, 1.20 [95% CI, 1.00 to 1.45]; n = 793; 3 studies) and functioning (Children's Global Assessment Scale pooled mean difference, 2.60 (95% CI, 0.78 to 4.42; n = 793; 3 studies). Other outcomes were not statistically significantly different. Differences in suicide-related outcomes and adverse events for pharmacotherapy when compared with placebo were not statistically significant. For suicide risk, no studies reported on the direct benefits of screening on health outcomes, and 2 RCTs (n = 2675) reported no harms of screening. One study (n = 581) reported on sensitivity of screening, ranging from 0.87 to 0.91; specificity was 0.60. Sixteen RCTs (n = 3034) reported on suicide risk interventions. Interventions were associated with lower scores for the Beck Hopelessness Scale (pooled mean difference, -2.35 [95% CI, -4.06 to -0.65]; n = 644; 4 RCTs). Findings for other suicide-related outcomes were mixed or not statistically significantly different.

CONCLUSION AND RELEVANCE Indirect evidence suggested that some screening instruments were reasonably accurate for detecting depression. Psychotherapy and pharmacotherapy were associated with some benefits and no statistically significant harms for depression, but the evidence was limited for suicide risk screening instruments and interventions.

JAMA. doi:10.1001/jama.2022.16310 Published online October 11, 2022.

- **Editorial**
- Multimedia
- Related article and JAMA Patient Page
- Supplemental content

Author Affiliations: RTI International-University of North Carolina at Chapel Hill **Evidence-based Practice Center** (Viswanathan, Wallace, Cook Middleton, Kennedy, Hudson, Rains, Kahwati); RTI International, Research Triangle Park, North Carolina (Viswanathan, Wallace, Kennedy, Hudson, Rains, Kahwati): Cecil G. Sheps Center for Health Services Research, University of North Carolina at Chapel Hill (Cook Middleton); Department of Psychiatry, University of North Carolina at Chapel Hill (McKeeman); Division of General Pediatrics and Adolescent Medicine, University of North Carolina at Chapel Hill (Vander Schaaf).

Corresponding Author: Meera Viswanathan, PhD, RTI International, 3040 E Cornwallis Rd, Research Triangle Park, NC 27709 (viswanathan@rti.org). epression and suicidal behaviors have long-term effects involving functional impairment, increased risk for substance abuse, and premature mortality. ¹⁻⁴ Routine screening may result in early identification and treatment. In 2016, the US Preventive Services Task Force (USPSTF) issued a recommendation for screening for major depressive disorder (MDD) in adolescents aged 12 to 18 years (B recommendation). ⁵ The USPSTF also concluded that the current evidence was insufficient to assess the balance of benefits and harms of screening in younger children. In 2014, the USPSTF concluded there was insufficient evidence to assess the balance of benefits and harms of screening for suicide risk in adolescents, adults, and older adults in primary care (I statement). ⁶ This review updated the evidence on screening for depression and suicide risk for children and adolescents to inform updated recommendations by the USPSTF.

Methods

Scope of the Review

The analytic framework and key questions (KQs) that guided the review are shown in Figure 1. Detailed methods, evidence tables, and contextual information are available in the full evidence report.⁸

Data Sources and Searches

PubMed, Cochrane Library, PsycINFO, CINAHL, and ClinicalTrials.gov were searched for English-language articles (eMethods in the Supplement). Depression searches were limited to articles published from January 1, 2015, through July 19, 2021; suicide risk searches were limited to articles published between June 1, 2012, and July 19, 2021. Evidence prior to these dates was identified from existing reviews. ^{9,10} Reference lists of pertinent articles and studies suggested by reviewers were also reviewed. Article alerts and targeted searches of journals to identify major studies published in the interim that may affect the conclusions or understanding of the evidence and the related USPSTF recommendation were used as part of ongoing surveillance. The last surveillance was conducted on June 1, 2022.

Study Selection

Two investigators independently reviewed titles, abstracts, and full-text articles using prespecified criteria for each KQ (eMethods in the Supplement); disagreements were resolved by discussion or by a third reviewer. English-language studies of persons 18 years or younger, on average, that met all study selection criteria, were fair or good methodological quality, and were conducted in countries categorized as very highly developed by the 2018 United Nations Human Development Index¹¹ were eligible. Studies included in the prior reviews for the USPSTF were reassessed against the study selection criteria. For screening, studies that included unselected participants without known risk of depression or increased risk of suicide were eligible. For depression, treatment studies that included at least half of participants with MDD were eligible. For suicide risk, treatment studies were restricted to participants with increased suicide risk. Eligible interventions included psychotherapy or first-line pharmacotherapy approved for pediatric use (eg, clonidine, duloxetine, fluoxetine, escitalopram, sertraline, fluvoxamine). Interventions were required to be relevant to or referable from primary care. Eligible outcomes for benefits

of screening and treatment included depression symptoms as measured through validated instruments, clinical response, or remission; suicide deaths, suicide attempts, and deliberate self-harm or suicidal ideation; all-cause mortality; quality of life measured using validated instruments; and functioning (validated scales, days of missed school, sleep-related outcomes). Eligible harms of treatment included treatment avoidance, deterioration in patient-clinician relationship, labeling or stigma, unnecessary treatment, serious adverse effects, withdrawals due to adverse effects, and suicidality.

Data Extraction and Quality Assessment

For each included study, 1 reviewer abstracted relevant study characteristics and outcomes into a structured form. A second reviewer checked all data for completeness and accuracy. Methodological quality ratings for studies included from a prior Agency for Healthcare Research and Quality (AHRQ) evidence review on depression treatment in youth 12 were spot-checked and carried forward. All other studies were assessed dually and independently using predefined criteria established by the USPSTF (eMethods in the Supplement) and others. 13-16

Disagreements in study quality ratings were resolved through discussion or by a third senior reviewer. Detailed study quality assessments are provided in eTables 1 through 7 in the Supplement.

Data Synthesis and Analysis

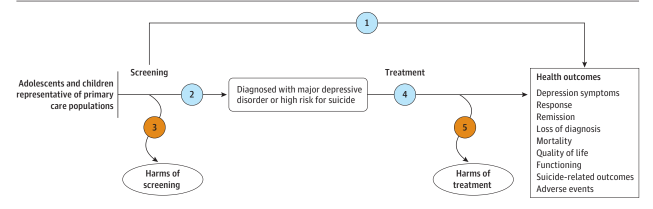
Data were synthesized in tabular and narrative forms. When at least 3 similar studies were available, a quantitative synthesis was performed using random-effects models with the inverse-variance weighted method of DerSimonian and Laird in Comprehensive Meta-Analysis version 3.3 to generate pooled estimates of effect.¹⁷ The *I*² statistic was calculated to assess statistical heterogeneity in effects.¹⁸ Significance testing was based on the exclusion of the null value by the 95% CI around the pooled estimate; all testing was 2-sided.

The strength of evidence was assessed as high, moderate, low, or insufficient using methods developed for the USPSTF and the AHRQ Evidence-based Practice Center program. Two senior reviewers independently developed initial strength-of-evidence assessments; disagreements were resolved through discussion or input of a third senior reviewer.

Results

Forty studies in 54 publications (N = 11220 from screening accuracy studies and randomized clinical trials [RCTs]) were eligible (Figure 2). For depression, 7 screening test accuracy studies $^{21-27}$ and 13 treatment RCTs $^{28-47}$ (N = 5433) were identified in addition to 1 meta-analysis. 48 Two studies of screening accuracy, 22,25 7 treatment trials, 28,32,33,35,37,38,40,42,47 and the meta-analysis 48 are new to this update. For suicide risk, 1 screening test accuracy study, 49 2 studies of screening harms, 50,51 and 16 treatment RCTs were identified (N = 6290). $^{52.74}$ Nine trials are new to this update. $^{52.55,58,60,64-72}$ The results in this publication focus on pooled analyses when available. Additional results are available in the full report. 8 A list of full-text articles that were screened but excluded is provided in the Supplement (List of Excluded Studies).

Figure 1. Analytic Framework: Screening for Depression or Suicide Risk in Children and Adolescents



Key questions

- Do depression or suicide risk screening programs in primary care or comparable settings result in improved health outcomes in children and adolescents?
- 2 Do instruments to screen for depression or suicide risk accurately identify children and adolescents with depression and increased risk of suicide in primary care or comparable settings?
- What are the harms associated with screening for depression or suicide risk in primary care or comparable settings in children and adolescents?
- 4 Does treatment (psychotherapy, pharmacotherapy, or collaborative care) of depression or suicide risk result in improved health outcomes in children and adolescents?
- What are the harms of treatment (psychotherapy, pharmacotherapy, or collaborative care) in children and adolescents who are treated for depression or suicide risk?

Evidence reviews for the US Preventive Services Task Force (USPSTF) use an analytic framework to visually display the key questions that the review addressed to allow the USPSTF to evaluate the effectiveness and safety of a

preventive service. The questions are depicted by linkages that relate interventions and outcomes. Refer to the USPSTF Procedure Manual for interpretation of the analytic framework.⁷ KQ indicates key question.

Benefits of Screening

Key Question 1. Do depression or suicide risk screening programs in primary care or comparable settings result in improved health outcomes in children and adolescents?

No trials directly assessing the benefits of screening children or adolescents in the primary care setting for MDD or suicide risk compared with no screening were found.

Accuracy of Screening Instruments

Key Question 2. Do instruments to screen for depression or suicide risk accurately identify children and adolescents with depression or increased risk of suicide in primary care or comparable settings?

Depression

Seven fair-quality studies of diagnostic test accuracy (n = 3316) were included $^{21-27}$; 2 were new to this update. $^{22.25}$ Authors assessed 7 different screening instruments (eTables 8 and 9 in the Supplement describe index tests and reference standards, respectively). Some authors assessed more than 1 instrument or more than 1 threshold for a positive screen result for the same instrument, and few studies prespecified thresholds for a positive screen result. All but 1 study 25 were restricted to adolescents.

The prevalence of MDD based on reference standard diagnostic clinical interviews ranged from 3% to 9% across studies enrolling persons recruited from school or community-based settings^{21,23,26,27} and was 11% in all 3 of the studies enrolling persons from nonpsychiatric clinical settings.^{22,24,25} **Table 1** provides a summary of sensitivity and specificity (additional detail is provided in eTable 10 in the Supplement). Excluding 2 outliers,²⁶ the sensitivity across these instruments for identifying MDD ranged from 0.73 to 0.94, and the specificity ranged from 0.38 to 0.97.

Suicide Risk

One fair-quality study (n = 580) conducted in the US that recruited potential high school dropouts aged 14 to 20 years evaluated the 20-item Suicide Risk Screen (SRS). 49 The prevalence of increased suicide risk ranged from 19% to 22%, depending on the reference standard used. The sensitivity and specificity of the SRS compared with one reference standard were 0.91 and 0.60, respectively, and were 0.87 and 0.60 against another reference standard. 49

Harms of Screening

Key Question 3. What are the harms associated with screening for depression or suicide risk in primary care or comparable settings in children and adolescents?

E3

jama.com JAMA Published online October 11, 2022

37706 Citations identified through database searcha 37157 Citations excluded at title and abstract review 549 Citations excluded after title and abstract review 249 Additional citations included 212 From previous review assessed at full-text stage 37 From hand searches and peer-review recommendations 798 Full-text articles assessed for eligibility for all KQs 744 Excluded 171 Population 115 Comparator 102 Intervention 78 Outcomes 59 Design 52 Other target condition 51 Quality 35 Setting 18 Publication type 17 Country 16 Protocol or ongoing study 15 Language 10 Duplicate **5** Superseded by publication **54** Articles (49 studies) included for depression and suicide risk topic O Articles included for KO1 2 Articles (2 studies) included 2 Articles (2 studies) included 43 Articles (29 studies) 13 Articles (9 studies) included for KQ4b included for KQ5^l for KQ2 for KQ3 6 Articles (4 studies) 2 Articles (2 studies) from 2 Articles (2 studies) from 18 Articles (13 studies) previous review previous review from previous review from previous review 0 Articles from current 0 Articles from current 25 Articles (16 studies) 7 Articles (5 studies) review from current review from current review

Figure 2. Literature Search Flow Diagram: Screening for Depression or Suicide Risk in Children and Adolescents

Reasons for exclusion: Population: Study was not conducted in an included population. Comparator: Study did not use an included comparator. Intervention: Study did not use an included intervention. Outcomes: Study did not report relevant outcomes. Design: Study did not use an included design. Other target condition: Study reported on anxiety. Quality: Study was poor quality. Setting: Study was not conducted in settings representative of primary care. Publication type: Publication was a commentary. Country: Study was not conducted in a country relevant to US practice. Protocol or ongoing study:

Study was a protocol or ongoing study and did not report eligible outcomes. Language: Study was not in English. Duplicate: Study was a duplicate of other studies in the review. Superseded by publication: Study findings were wholly superseded by another publication.

Depression

No trials directly assessing the harms of screening children or adolescents in the primary care setting for MDD compared with no screening were found.

Suicide Risk

Two fair-quality RCTs (n = 2675) conducted in high schools reported no differences in harms in distress, transient mood states, or suicidal ideation after screening for suicide risk. 50,51

Benefits of Treatment

Key Question 4. Does treatment (psychotherapy, pharmacotherapy, or collaborative care) of depression or suicide risk result in improved health outcomes in children and adolescents?

Benefits of treatment for depression and increased suicide risk are summarized in **Table 2**.

Depression

Thirteen fair-quality RCTs (20 publications) were identified (n = 2152). ²⁸⁻⁴⁷ Seven RCTs ^{28,32,33,35,37,38,40,42,47} were new in this update. Key characteristics of included depression studies are provided in eTable 11 in the Supplement, detailed outcomes are provided in eTables 12 through 27 in the Supplement, and results from meta-analyses are provided in eFigures 1 through 8 in the Supplement.

Eleven RCTs enrolled youth meeting *Diagnostic and Statistical Manual (DSM)* criteria for MDD.^{28,29,31,33-40} Two RCTs enrolled a sample in which more than 50% of participants met *DSM* criteria for MDD but also admitted youth with MDD, dysthymia, or depressive disorder not otherwise specified.^{30,32} Mean ages ranged from 5 to 17.5 years.^{33,38,42} One RCT focused on early childhood (3-6 years),^{33,42} 2 focused on school-aged children and adolescents (ages ranged from 7-14 years and 6-17 years),^{32,39} and

JAMA Published online October 11, 2022

^a Combined searches were conducted on anxiety, depression, and suicide risk. Results for anxiety are presented in a separate publication.²⁰

^b Study may address more than 1 key question (KQ).

Table 1. Accuracy of Screening Instruments for Screening for Depression and Suicide Risk

Condition	Screener	Age range	No. of studies	Sensitivity	Specificity
Major depressive	Beck Depression Inventory (threshold = 11)	15-18.5	2 ^{21,27}	0.84-0.90	0.81-0.86
disorder	Center for Epidemiologic Studies Depression Scale (various thresholds)	12-18	2 ²⁶	0.18 ^a -0.85	0.38-0.83
	Clinical Interview Schedule-Revised	Mean, 15.7	1 ²⁶	0.18 ^a	0.97 ^b
	Hopkins Symptom Checklist	14-16	1 ²²	0.85	0.78
	Patient Health Questionnaire-Adolescents	13-18	1 ²⁴	0.73	0.94
	Paediatric Index of Emotional Distress- Depression subscale	8-17	1 ²⁵	0.94	0.81
	World Health Organization 5-Item Well-being Index	14-16	1 ²²	0.88	0.80
Suicide risk	Suicide Risk Screen	14-20	1 ⁴⁹	0.87-0.91 ^c	0.60

- ^a Across the 2 studies, 5 thresholds were evaluated. The sensitivity for most ranged between 0.59 and 0.85 except for the thresholds of 20 or greater and 22 or greater among boys, which were 0.19 and 0.18, respectively.
- ^b These estimates were from an analysis weighted for selection into the second phase of the study; calculated sensitivity without weighting was 0.74 and specificity was 0.78.
- ^c This instrument was evaluated against 2 different reference standards.

10 focused on adolescents (ages ranged from 12-17 years and 15-19 years). ^{28-31,34-38,40} Two had majority male participants. ^{32,33,42}

Two pharmacotherapy trials compared escitalopram with placebo. 31,39 One 3-group trial compared fluoxetine, cognitive behavioral therapy (CBT), and placebo. 34 One trial compared collaborative care with enhanced usual care. 36 Six studies focused on CBT, 28-30,32,37,38 and 3 focused on counseling other than CBT. 33,35,40 Comparisons included treatment as usual, wait-list, placebo, attention control, and supportive contact. The results reported here focus on psychotherapy and pharmacotherapy trials; detailed results for collaborative care and combination therapy are available in the full report. 8

Outcomes reported included (1) depression symptoms; (2) response, remission, or loss of diagnosis; and (3) quality of life and functioning. Regarding depression symptoms, pooled estimates suggested improved symptoms associated with psychotherapy (Beck Depression Inventory [BDI] pooled standardized mean difference, -0.58 [95% CI, -0.83 to -0.34]; n = 471; 4 RCTs^{30,35,37,38} and Hamilton Depression Scale pooled mean difference, -2.25 [95% CI, -4.09 to -0.41]; n = 262; 3 RCTs^{29,30,35}) or pharmacotherapy (Children's Depression Rating Scale-Revised mean difference, -3.76 [95% CI, -5.95 to -1.57]; n = 793; 3 RCTs^{31,34,39}) in some but not all measures of symptom severity. Regarding clinical response, 3 studies reported responses on the BDI and BDI-II (BDI version 2) scale^{35,37,38} favoring pharmacotherapy but results could not be pooled because of the various thresholds used. Results favored pharmacotherapy for remission (relative risk [RR], 1.20 [95% CI, 1.00 to 1.45]; n = 793; 3 RCTs^{31,34,39,45}) and psychotherapy for loss of diagnosis (RR, 1.73 [95% CI, 1.00 to 3.00]; n = 395; 4 RCTs^{30,34,37,38}), but CIs included the null. Studies reported improved functioning, measured by the Children's Global Assessment Scale for pharmacotherapy (mean difference, 2.60 [95% CI, 0.78 to 4.42]; n = 793; 3 RCTs $^{31,34,39})$ but not for psychotherapy (mean difference, 1.52 [95% Cl. -1.54 to 4.58l: n = 601: 4 RCTs). 28,29,34,35

Two psychotherapy studies focused on or included children; 1 reported improvements in symptoms, loss of diagnosis, and functioning, ³³ and a second reported no differences for symptoms or remission. ³² One pharmacotherapy study that included children reported no differences in symptoms, response, or functioning. ³⁹

Suicide Risk

Sixteen RCTs of good or fair quality (described in 23 articles) were identified. $^{52.74}$ Nine were new to this update (n = 3034). $^{52.55,58.60,64.72}$

Key characteristics of included studies are described in eTable 28 in the Supplement, detailed outcomes included in pooled results are provided in eTables 29 and 30 in the Supplement, and results from meta-analyses are provided in eFigures 9 through 15 in the Supplement.

Fourteen studies enrolled adolescents based on elevated suicide risk, $^{52-55,57\cdot72,74}$ and 2 studies admitted adolescents with suicide risk and self-reported depressive symptoms (BDI $>19^{73}$ or BDI $>20^{56}$). Mean ages ranged from 14 to 18 years. All included studies focused on adolescents $^{52-74}$ and included a majority of female participants.

All included studies examined psychotherapy, counseling, support, or a combination with variable intensity and duration. Fifteen studies compared these interventions with treatment as usual, 52-59,61-74 which often included active psychosocial treatment, and 1 study compared the intervention with attention control. 60 Duration of treatment ranged between 1 single session to weekly sessions over 12 months. No evidence was captured that examined pharmacotherapies.

Outcomes reported included (1) suicide deaths, (2) suicide attempts and deliberate self-harm or suicidal ideation, (3) all-cause mortality, and (4) functioning. Three studies reported no deaths or no statistically significant differences in suicide deaths at the end of treatment (19 weeks to 12 months). 57,62,65-68 Pooled results for suicide attempt, deliberate self-harm, or suicidal ideation generally indicated no statistically significant differences for self-harm events (RR of self-harm events, 0.88 [95% CI, 0.63 to 1.24]; n = 1040; 5 RCTs $^{53-55,57,59,72,74}$; mean difference in self-harm events, -0.76 [95% CI, -2.15 to 0.63]; n = 972; 3 RCTs^{53-55,65-68,74}) or suicide-related hospitalization or emergency use (RR, 1.00 [95% CI, 0.67 to 1.50]; n = 978; 3 RCTs^{53-55,65-70}). The exception was measures of suicidal ideation: studies reported benefits associated with the intervention on the Beck Hopelessness Scale (mean difference, -2.35 [95% CI, -4.06 to -0.65]; n = 644; 4 RCTs^{62,64-68,73}). Pooled results for the Suicidal Ideation Questionnaire (SIQ) and SIQ-Junior scales also favored the intervention group but were not statistically significantly different (standardized mean difference, -0.18 [95% CI, -0.36 to 0.01]; n = 1111; 7 RCTs^{56,57,59,62,64-68,74}). Regarding all-cause mortality, a long-term follow-up of a study on a youth-nominated support team approach, 62 11 to 14 years after psychiatric hospitalization for suicide risk (baseline for the study), found a higher number of deaths in the National Death Index in the treatment-as-usual group when compared with the active treatment group (13/225 vs 2/223; hazard ratio, 6.62 [95% CI, 1.49 to 29.35]). 63 The same study did not

	⋧	
	-	ī
	₹	
		ū
	t	ä
	_	۰
	₹	
	c	١
	۲	ī
`		

Table 2. Benefits for Treatment for Screening for Depression and Suici	tment for Screen	ing for Depression ar	ıd Suicide Risk					
Intervention	Time of outcome measurement	Outcome measure, range, threshold	Outcome range	Outcome threshold indicating clinically meaningful effect	Treatment range at follow-up	Comparator range at follow-up	No. of studies (No. of participants)	Pooled results (95% CI) ^a ; I ²
Depression—change in symptoms	ptoms							
Internet-based individual CBT group; in-person CBT with and without parents; interpersonal psychottherapy ^b	8-12 wk	BDI or BDI-II	BDI: 0-63 ⁷⁵	BDI: Score <10: minimal depression Score 10-18: mild to moderate depression Score 19-29: moderate to severe depression Score ≥30: severe depression ^{21,27} BDI-II: Score 0-13: minimal depression Score 14-19: mild depression Score 20-28: moderate depression Score 20-28: severe depression	BDI-II: 16-19.9	BDI: 12.3-16 BDI-II: 24.8-25.2	4 30,35,37,38	Standardized mean difference, -0.58 (-0.83 to -0.34) l² = 0%
Individual in-person youth CBT; group in-person CBT with and without parents; interpersonal psychotherapy ^b	8-52 wk from baseline	НАМ-D	Unclear (2 studies used a 14-item version of HAM-D)	Unclear	4.9-8.7	6.5-12.8	3 ^{29,30,35} (262)	Mean difference, -2.25 (-4.09 to -0.41) 1 ² = 0%
Individual in-person CBT; family CBT	12-52 wk from baseline	CDRS-R	17-113	Score ≥40 indicates depression Score ≤28 indicates remission (minimal or no symptoms)	30.0-42.1	28.2-41.8	3 ^{28,32,34} (471)	Mean difference, 0.77 (-0.97 to 2.48) $I^2 = 0\%$
Escitalopram; fluoxetine	8-12 wk from baseline	CDRS-R	17-113	Score ≥40 indicates depression Score ≤28 indicates remission (minimal or no symptoms)		36.4-41.8	3 ^{31,34,39} (793)	Mean difference, -3.76 (-5.95 to -1.57) 1 ² = 49%
Depression—remission and loss of diagnosis	loss of diagnosis							
CBT	8-12 wk from baseline	Loss of diagnosis measured by clinical interviews	0%-100% For proportion	NA	56%-71%	16%-60%	4 ^{30,34,37,38} (395)	RR, 1.73 (1.00 to 3.00) $I^2 = 81\%$
Escitalopram; fluoxetine	8-12 wk from baseline	Remission from depression symptoms (CDRS-R≤28)	0%-100% For proportion	CDRS-R < 28 indicates moderate marked improvement, proportion threshold unclear	23%-46%	17%-38%	3 ^{31,34,39,45} (793)	RR, 1.20 (1.00 to 1.45) <i>I</i> ² = 0%
Depression—change in functioning	tioning							
Individual in-person CBT; interpersonal psychotherapy	12-52 wk from baseline	CGAS	1-100	Score >70: no clinically significant functional impairment Score <41: major impairment to functioning in several areas	60.0-72.3	59.3-74.1	4 ^{28,29,34,35} (601)	Mean difference, 1.52 (-1.54 to 4.58) $I^2 = 66\%$
Escitalopram; fluoxetine	8-12 wk from baseline	CGAS	1-100	Score >70: no clinically significant functional impairment Score <41: major impairment to functioning in several areas	62.1-68.5	59.3-64.6	3 ^{31,34,39} (793)	Mean difference, 2.60 (0.78 to 4.42) $I^2 = 0\%$

lable 2. Benefits for Treatment for Screening for Depression and Suicide Kisk (continued)	tment for screer	ııng ror Depression al	o suicide Kisk (co	ntinued)				
Intervention	Time of outcome measurement	Outcome measure, range, threshold	Outcome range	Outcome threshold indicating clinically meaningful effect	Treatment range at follow-up	Comparator range at follow-up	No. of studies (No. of participants)	No. of studies (No. of participants) Pooled results (95% CI) ^a ; <i>I</i> ²
Suicide risk-suicide-related outcomes	d outcomes							
Group psychotherapy; family therapy; mentalization-based treatment; developmental group therapy	6-18 mo	Proportion with self-harm events	0-100	NA	0.55%-88%	1.1%-83%	553-55,57,59,72,74	RR, 0.88 (0.63 to 1.24) I ² = 80%
Family therapy; DBT; developmental group therapy	19 wk to 18 mo	Mean No. of self-harm events	NA	NA	0.6-9.0	1.2-22.50	3 ⁵³⁻⁵⁵ ,65-68,74 (972)	Mean difference, -0.76 (-2.15 to 0.63) $I^2 = 68\%$
Youth-nominated support team; motivational interviewing; DBT; IPT-A-IN	2 mo to 19 wk	Suicidal ideation: BHS	0-20	Score > 9 indicative of suicide intentions	5.66-7.74	7.80-12.42	4 ^{62,64-68,73} (644)	Mean difference, -2.35 (-4.06 to -0.65) $I^2 = 46\%$
Attachment-based family therapy; group psychotherapy; group therapy; group therapy youth-nominated support team; motivational interviewing; DBT; developmental group therapy	2 mo to 71 wk	SIQ or SIQ-JR	SIQ: 0-180 SIQ-JR: 0-90	SIQ score >41 indicative of suicidal ideation SIQ-IR score >31 indicative of suicidal ideation	SIQ: 41.3-74.11 SIQ-JR: 5.2-25.55	SIQ: 39.7-76.40 SIQ-JR: 16.2-29.71	756.57,59.62,64-68,74 (1111)	SMD, -0.18 (-0.36 to 0.01) 1 ² = 45%
Family therapy; DBT; therapeutic assessment	18 mo to 2 y	Proportion with suicide-related hospitalization or emergency department use	NA	NA	1%-88%	1%-94%	3 53-55,65-70 (978)	RR, 1.00 (0.67 to 1.50) 1 ² = 21%
Attachment-based therapy and internet CBT	8-24 wk	Clinical response: SIQ-JR or perceived burdensomeness	SIQ-JR: 0-90; Perceived burden- someness: 6-42	Clinical response: SIQ-JR scores (defined as <13) Perceived burdensomeness <14.61	SIQ-JR: 87% Perceived burden- someness: 24%	SIQ-JR: 52% Perceived burden- someness: 10%	2 ^{56,60} (146)	Results from individual studies: OR for SIQ-JR clinical response, 6.30 (1.76-22.61) ⁵⁶ No significant differences at 24 wk for perceived burdensomeness (OR, 2.82 [0.80-9.91]) ⁶⁰
Suicide risk—change in functioning	ctioning							
Group psychotherapy; group therapy; developmental group thrapy; psychoeduca-tion for parents	8 wk to 7 mo	Functioning: Health of the Nation Outcome Scales for Children and Adolescents	0-52	Scores >13 indicate impairment of clinical significance	8.4-16.8	6.9-17.6	457,59,71,74 (509)	Mean difference, -0.40 (-2.55 to 1.78) $I^2 = 56\%$
Therapeutic assessment; individual and family DBT; group therapy	8-71 wk	Functioning: CGAS	1-100	Score >70: no clinically significant functional impairment Score <41: major impairment to functioning in several areas	58.5-65.7	60.1-64.22	3 ^{59,66,70} (195)	Mean difference, 1.30 (-2.52 to 5.12) $I^2 = 30\%$
Abbreviations: BDI, Beck Depression Inventory: BDI-II, Beck Depression Inventory version 2: Beck Hopelessness Scale.	nression Inventory	· BDI-II Beck Denression	Inventory version		a Results from pooled analyses unless otherwise specified	s unless otherwise	e specified	

Abbreviations: BU, Beck Depression Inventory; BU:III, Beck Depression Inventory version 2: Beck Hopelessness Scale, CBT, cognitive behavioral therapy; CDRS-R, Children's Depression Rating Scale-Revised; CGAS, Children's Global Assessment Scale; DBT, dialectical behavior therapy; HAM-D, Hamilton Depression Rating Scale; IPT-A-IN, intensive interpersonal bxpsychotherapy for depressed adolescents with suidal Irisk; NA, not applicable; OR, odds ratio; RR, relative risk; SIQ, Suicidal Ideation Questionnaire; SIQ-JR, Suicidal Ideation Questionnaire-Junior; SMD, standardized mean difference.

 $^{\rm b}$ The results across groups for the study were averaged (mean values calculated), with multiple treatment groups (group in-person CBT with or without parents³⁰) compared with wait-list.

© 2022 American Medical Association. All rights reserved.

demonstrate an effect on the primary outcome of suicidal ideation; as a result, findings by chance or through other mechanisms of action (such as improved problem solving) cannot be ruled out. The studies reported no statistically significant differences in functioning on the Health of the Nation Outcome Scales for Children and Adolescents (mean difference, -0.40 [95% CI, -2.55 to 1.78]; n = 509; 4 RCTs^{57,59,71,74}) or the Children's Global Assessment Scale (mean difference, 1.30 [95% CI, -2.52 to 5.12]; n = 195; 3 RCTs^{59,66,70}).

Harms of Treatment

Key Question 5. What are the harms of treatment (psychotherapy, pharmacotherapy, or collaborative care) in children and adolescents who are treated for depression or suicide risk?

Depression

Seven studies (described in 12 articles) were included (n = 1408 from primary studies). 28,31,34,36,39-41,43-46,48 All 7 studies are also included in the discussion of benefits except for 1 meta-analysis, which was new to this update. 48 Key characteristics of included depression studies are provided in eTable 31 in the Supplement, and detailed outcomes are provided in eTables 32 through 40 in the Supplement. Studies reported on suicide-related events (suicide deaths, attempts, deliberate self-harm, or suicidal ideation) and other adverse events. Regarding psychotherapy, the Treatment of Adolescents With Depression (TADS) trial with 2 active interventions (CBT and fluoxetine) reported varying results in the placebo group in different publications; as a result, the relative risk of suicide-related outcomes could not be calculated with certainty. A second trial reported 5 of 106 (4.7%) suicide-related events for CBT plus treatment as usual vs 2 of 106 (1.9%) for treatment as usual alone²⁸; differences were not statistically significant. For pharmacotherapy, inconsistent reporting across different publications on harms from the TADS trial led to uncertainty regarding the relative risk of suiciderelated outcomes with fluoxetine vs placebo. Other evidence for escitalopram studies indicated rates of suicide-related outcomes that are not statistically significantly different compared with placebo $(1/129 [0.8\%] \text{ vs } 2/132 [1.5\%]^{39}; 6/57 [3.8\%] \text{ vs } 6/155 [3.9\%]^{31}). \text{ The}$ meta-analysis also reported similar rates for escitalopram vs placebo (15/290 [5%] vs 15/294 [5%]; 2 studies) and fluoxetine vs placebo (51/521 [10%] vs 44/514 [9%]; 7 studies). 48 No statistically significant differences were reported for other harms for psychotherapy (deteriorated on the Quick Inventory of Depressive Symptomatology for Adolescents: 1 vs 3⁴⁰; harm-related adverse events: 5/111 participants [4.5%] vs 6/112 participants [5.4%]; OR, O.80 [95% CI, O.25 to 2.81]³⁴). For pharmacotherapy, 2 escitalopram trials reported no statistically significant differences for withdrawals (4/155 [2.6%] vs 1/157 [0.6%]³¹; 2/131 [1.5%] vs 2/133 [1.5%]³⁹) or serious adverse events (4/155 [2.6%] vs 2/157 [1.3%]³¹; 2/131 [1.5%] vs 3/133 [2.3%]³⁹). The TADS trial reported a higher but not statistically significant difference in adverse events for fluoxetine compared with placebo (13/109 participants [11.9%] vs 6/112 [5.4%]; OR, 2.4 [95% CI, 0.87 to 6.54]).34

Suicide Risk

Two studies^{53-55,58} reported on adverse events (n = 885). One study⁵³⁻⁵⁵ reported on adverse events, serious adverse events, and other harms during the 12- to 18-month follow-up period. Similar numbers of adverse events, including attendance at minor injury

units, walk-in centers, and accident and emergency centers and rereferral to mental health services, occurred in the family therapy group (54%) and treatment-as-usual group (52%). Serious adverse events, defined as hospital attendance, also occurred at similar rates across the intervention (38%) and control (34%) groups. Two participants assigned to the family therapy group died between 3 and 4 years after randomization. Neither death was related to self-harm. One additional study⁵⁸ reported 5 adverse events among 4 participants, but the occurrences were not considered trial-related and were not reported by group.

Discussion

This systematic review evaluated screening for depression and suicide risk in children and adolescents. **Table 3** summarizes the evidence, including strength-of-evidence ratings. No studies reported on the direct benefits or harms of screening. The discussion below focuses on the indirect evidence from studies describing test accuracy, benefits of treatment, and harms of treatment.

Depression

The standard of evidence for test accuracy was rated as low to moderate for sensitivity and moderate for specificity.

The depression module (Patient Health Questionnaire 9 [PHQ-9]) of the full PHQ is the instrument highlighted for use in screening for depression by the American Academy of Pediatrics. 6 One study of the accuracy of the full PHQ modified for adolescents was included, but no studies evaluating the PHQ-9 were identified. Based on the accuracy characteristics for the 1 included study of the Patient Health Questionnaire—Adolescents (PHQ-A), 24 per 1000 screening tests conducted, 58 false-positives and 8 falsenegatives would be generated at the low end of MDD prevalence (3%) observed in this updated review, and 53 false-positives and 30 false-negatives would be generated at the high end of prevalence (11%). Positive screening results would require additional diagnostic evaluation to determine true-positives from false-positives, but it is likely that some youth screening positive but not meeting diagnostic criteria for MDD may have persistent depressive disorder (formerly known as dysthymia) or other behavioral health conditions with symptoms similar to depression. The consequences of a false-negative would largely depend on the severity of the missed diagnosis; the likelihood of missing a severely depressed youth is small because most screen-detected depression is likely to be mild to moderate. However, even mildly to moderately depressed youth may have suicidal ideation, and the consequences of missing such symptoms could be serious.

The updated evidence on psychotherapy suggested some benefits for symptom improvement and clinical response, but the results were not consistent across all measures for other outcomes. The evidence for pharmacotherapy suggested benefit for symptom improvement, but the results were not consistent across all measures for other outcomes. Thus, the strength of evidence for psychotherapy and pharmacotherapy was rated as low for benefit. The evidence on harms was very limited; no statistically significant differences were observed. One multigroup trial (TADS) with inconsistent reporting on suicide-related events across its various publications contributed to the evidence on psychotherapy,

Table 3. Summary of Evidence					
No. of studies, study designs (No. of participants)	Summary of findings	Consistency and precision	Limitations	Strength of evidence	Applicability
KQ1: Benefits of screening					
None	NA	NA	NA	Insufficient	NA
KQ2: Accuracy of screening					
Depression: 7 studies (n = 3281) ²¹⁻²⁷	Varies by screener and threshold, excluding outliers Sensitivity range, 0.59-0.94 Specificity range, 0.38-0.96 PHQ-A: Sensitivity, 0.73 (95% CI, 0.58-0.85) Specificity, 0.94 (95% CI, 0.91-0.96)	Consistent when multiple studies are available, precise for specificity, precision varies for sensitivity	Unclear whether thresholds were established a priori or whether index and reference standard results were blinded; no replication of approaches for most screeners	Low to moderate for sensitivity (varies by instrument); moderate for specificity	Primarily adolescents, as only 1 study included children younger than 12 y; 7 different screeners evaluated but most not being used in practice
Suicide: 1 study (n = 581)	Sensitivity, 0.87 and 0.91 (varies by reference standard) Specificity, 0.60	Consistency unknown; imprecise	Unclear whether thresholds were established a priori or whether interviewers were blinded; single study	Insufficient	Participants were potential high school dropouts; instrument was a 20-item screener embedded into a longer questionnaire, so unclear whether feasible in primary care
KQ3: Harms of screening tests					
Depression: none	NA	NA	NA	Insufficient	NA
Suicide: 2 RCTs ^{50, 51} (n = 2675)	No significant difference in suicidal ideation between students exposed to screening items and those not exposed (1 RCT) No significant differences in measures of short-term distress/emotions for students exposed to suicide screening items compared with those not exposed (2 RCTs)	Consistent; precise	Fair-quality trials with some attrition; only evaluated measures of immediate and short-term emotions (over 1-2 d)	Low for no short-term harms from screening for suicide risk; insufficient for screening for depression and anxiety	High school students; 1 study entirely comprised boys
KQ4: Benefits of treatment					
Depression: 13 RCTs ^{28-40,42} (2 on pharmacotherapy; 9 on psychotherapy; 1 on CBT, fluoxetine, and their combination; 1 on collaborative care) (n = 2152)	Psychotherapy Varied by measure, with some pooled estimates of effect favoring psychotherapy for symptoms, clinical response, and loss of diagnosis, but other outcome measures did not consistently demonstrate a statistically significant difference Pharmacotherapy: Statistically significant differences favoring pharmacotherapy for 1 measure of symptoms Pooled differences favored pharmacotherapy but were not statistically significant difference Other outcome measures did not demonstrate a statistically significant difference Collaborative care: Statistically significant differences favoring collaborative care for symptoms at 6 mo and clinical response by 12 mo; remission at 6 mo; no benefits for functioning	Mostly consistent; mostly imprecise	Psychotherapy cannot mask treatment, leading to the potential for bias in outcome reporting	Psychotherapy: low for benefit for all outcomes other than remission Pharmacotherapy: low for benefit for all outcomes other than response Collaborative care: low for benefit for symptoms, response, and remission; insufficient for functioning	Studies addressed youth aged 3 to 19 y, but 9 were conducted exclusively in adolescents Pharmacotherapy studies were limited to first-line drugs with FDA approval for pediatric use

Table 3. Summary of Evidence (continued)	(pai				
No. of studies, study designs (No. of participants)	Summary of findings	Consistency and precision	Limitations	Strength of evidence	Applicability
Suicide: 16 RCTs ^{52-62,64-74} (3034)	Statistically significant difference favoring interventions for suicidal ideation on the Beck Hopelessness Scale, nonstatistically differences favoring suicide risk interventions on the SiQ and SiQ-JR, mixed on other measures No statistically significant differences on suicide deaths, hospitalization or emergency department visits, number of self-harm events, proportion with self-harm events, or functioning All-cause mortality: statistically significant difference favoring interventions (hazard ratio for treatment as usual, 6.62 [95% CI, 1.49-29.35]; n = 448; 1 study)	Consistent; imprecise	mask treatment, leading to the leading to the potential for bias in outcome reporting, all comparison groups are treatment-as-usual comparisons, which in many cases were active treatments and could bias results toward null effects	Psychotherapy: Low for benefit for suicidal ideation and clinical response; insufficient for all other outcomes	Applicable to adolescents (predominantly females); no studies recruited children younger than 11 y; most recruited from mental health or specialist settings
KQ5: Harms of treatment					
Depression: 6 RCTs ^{28,31,34,36,39-41,43-46} (3 on pharmacotherapy; 2 on psychotherapy; 1 on CBT, fluoxetine, and their combination; 1 on collaborative care) (n = 1408 from trials) and 1 meta-analysis ⁴⁸	Psychotherapy: no statistically differences in negative effects in 2 trials; however, precise effects unclear because 1 trial had inconsistent reporting. Pharmacotherapy: nonsignificant but higher risk of suicide-related outcomes, withdrawal due to adverse events and serious adverse events, precise effects unclear owing to inconsistent study reporting. Collaborative care: inconsistent results for psychiatric hospitalizations and emergency department visits	Consistent to inconsistent; imprecise	Psychotherapy trials cannot mask treatment, leading to the potential for bias in outcome reporting, inconsistent results across publications from 1 trial	Psychotherapy: insufficient Pharmacotherapy: low for harms Collaborative care: insufficient	Studies addressed youth aged 6 to 18 y, but 5 were conducted exclusively in adolescents Pharmacotherapy studies were limited to first-line drugs with FDA approval for pediatric use
Suicide: 2 RCTs ^{33, 58} (885)	No statistically significant differences on adverse events (such as minor injury, walk-in, accident and emergency centers, re-referral to mental health service, and hospital attendance)	Consistent; imprecise	mask treatment, leading to the potential for bias in outcome reporting, all comparison groups are treatment-as-usual comparisons, which in many cases were active treatments and could lead to bias stoward null effects	Insufficient	Applicable to adolescents, primarily females, recruited from mental health or specialist settings
Abbreviations: CBT cognitive behavioral th	Abbreviations: CBT cognitive behavioral therany: FDA 11S Food and Drug Administration: KO kev nuestion. Na. not applicable: PHO:A Patient Health Questionnaire-Adolescents: RCT randomized clinical trial: SIO. Suicidal Ideation	setion. NA not applicable. F	HO-A Patient Health Our	estionnaire-Adolescents-RCT r	andomized clinical trial: SIO Suicidal Ideation

Abbreviations: CBT, cognitive behavioral therapy; FDA, US Food and Drug / Questionnaire; SIQ-JR, Suicidal Ideation Questionnaire-Junior.

JAMA Published online October 11, 2022 jama.com

E10

pharmacotherapy, and their combination. These discrepancies increased the uncertainty regarding harms of treatment and have led to a call for independent reanalysis of the TADS results. ^{77,78} The US Food and Drug Administration (FDA) noted a higher frequency of suicide-related events in boxed warnings for antidepressants. ⁷⁹ The underlying FDA review for this warning relied on drug trials in populations ineligible for this review. ⁸⁰

Suicide

Only 1 eligible study assessed the accuracy of screening for suicide risk in adolescents evaluated against a clinical diagnostic interview reference standard; the instrument used was the SRS, a 20-item instrument embedded in a longer questionnaire, and the study population was recruited from youth identified as potential high school dropouts. 49 The strength of evidence for screening was rated as insufficient because of inconsistencies in estimates based on the reference standard used, imprecision, and study limitations. Given that many depression screening instruments include an assessment of suicidal ideation, it is unclear whether a separate, stand-alone instrument to screen for increased suicide risk has value for universal screening in primary care practice. The Ask Suicide Screening Questions (ASQ) is a brief 4-item instrument that was initially developed for youth 8 years or older in emergency department settings but has since been evaluated in other medical settings, including outpatient specialty and primary care. 81,82 The Joint Commission recommends suicide risk screening for all medical patients in all medical settings, including outpatient practices.⁸³ The National Institute for Mental Health developed an ASQ toolkit to support implementation of suicide risk screening in medical settings, including for youth in primary care.⁸⁴ One study evaluating the ASQ in outpatient settings, including primary care, was identified, but it was excluded because its accuracy was compared against another suicide risk screening instrument and not against a diagnostic clinical interview by a qualified professional.⁸²

The updated evidence base suggests improvements in suicidal ideation resulting from psychotherapy interventions, but this finding was statistically significant for only 1 measure. The evidence suggested no statistically significant differences on all other measures. All trials included treatment-as-usual comparators, which for ethical reasons must be active comparators, such as standard psychotherapy, individual counseling, family sessions, medication assessment and review, medication, and other care coordination activities. Comparable intensity of therapy in study groups, coupled with low event rates for some outcomes (such as suicide deaths, hospitalizations, and suicide attempts), is likely to make differences between study groups difficult to detect. The evidence was rated as low for benefit on suicidal ideation; insufficient for evaluating outcomes such as suicide attempts, hospitalizations, and deaths; and low for no harm.

Limitations

This study has several limitations. First, no studies were available that compared screening with no screening. Second, only limited evidence was available on long-term outcomes, test accuracy, and suicide risk and depression treatment in children. Third, treatment-asusual comparators for suicide risk interventions included active treatments, which may have led to lack of statistically significant differences between study groups. Fourth, the review was limited to first-line drugs approved for pediatric use by the FDA.

Conclusions

Indirect evidence suggested that some screening instruments were reasonably accurate for detecting depression. Psychotherapy and pharmacotherapy were associated with some benefits and no statistically significant harms for depression, but the evidence was limited for suicide risk screening instruments and interventions.

ARTICLE INFORMATION

Accepted for Publication: August 25, 2022.

Published Online: October 11, 2022. doi:10.1001/jama.2022.16310

Author Contributions: Dr Viswanathan had full access to all of the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

Concept and design: Viswanathan, Kahwati.
Acquisition, analysis, or interpretation of data:
Viswanathan, Wallace, Cook Middleton, Kennedy,
McKeeman, Hudson, Rains, Vander Schaaf.
Drafting of the manuscript: Viswanathan, Kennedy,
Rains.

Critical revision of the manuscript for important intellectual content: All authors.

Statistical analysis: Viswanathan, Wallace, Kahwati.
Obtained funding: Viswanathan, Kahwati.
Administrative, technical, or material support:
Viswanathan, Kennedy, Rains.
Supervision: Viswanathan.

Conflict of Interest Disclosures: Dr Vander Schaaf reported that she is a Fellow of the American Academy of Pediatrics and member of American Academy of Pediatrics CATCH Committee. No other disclosures were reported.

Funding/Support: This study was funded under contract HHSA-290-2015-00011-I, Task Order 15, from the Agency for Healthcare Research and Quality (AHRQ), US Department of Health and Human Services (HHS), under a contract to support the US Preventive Services Task Force (USPSTF).

Role of the Funder/Sponsor: Investigators worked with USPSTF members and AHRQ staff to develop the scope, analytic framework, and key questions for this review. AHRQ had no role in study selection, quality assessment, or synthesis. AHRQ staff provided project oversight, reviewed the report to ensure that the analysis met methodological standards, and distributed the draft for peer review. Otherwise, AHRQ had no role in the conduct of the study; collection, management, analysis, and interpretation of the data; or preparation, review, or approval of the manuscript findings. The opinions expressed in this document are those of the authors and do not represent the official position of AHRQ or HHS.

Additional Contributions: We acknowledge the following individuals for their contributions to this project: AHRQ staff Iris Mabry-Hernandez, MD, MPH, and Tracy Wolf, MD, MPH; current and former members of the USPSTF who contributed to topic deliberations; RTI International–University of North Carolina Evidence-based Practice Center staff

Christiane Voisin, MSLS, Candi Wines, MPH, Nila A. Sathe, MA, MLIS, Carol Woodell, BSPH, Sharon Barrell, MA, Staci Rachman, BA, Michelle Bogus, Sarah Barringer, and Loraine Monroe. USPSTF members, peer reviewers, and federal partner reviewers did not receive financial compensation for their contributions.

Additional Information: A draft version of the full evidence report underwent external peer review from 4 content experts (Gregory Simon, MD, MPH, University of Washington; Joan Asarnow, PhD, University of California, Los Angeles; Natalie Cort, PhD, William James College; and Raquel Halfond, PhD, American Psychological Association) and a federal partner reviewer (Erin Abramsohn, DrPH, MPH, Centers for Disease Control and Prevention). Comments from reviewers were presented to the USPSTF during its deliberation of the evidence and were considered in preparing the final evidence review.

Editorial Disclaimer: This evidence report is presented as a document in support of the accompanying USPSTF recommendation statement. It did not undergo additional peer review after submission to *JAMA*.

REFERENCES

- 1. Wilson S, Hicks BM, Foster KT, McGue M, lacono WG. Age of onset and course of major depressive disorder: associations with psychosocial functioning outcomes in adulthood. *Psychol Med*. 2015;45(3):505-514. doi:10.1017/
- 2. Whalen DJ, Dixon-Gordon K, Belden AC, Barch D, Luby JL. Correlates and consequences of suicidal cognitions and behaviors in children ages 3 to 7 years. *J Am Acad Child Adolesc Psychiatry*. 2015;54 (11):926-37. doi:10.1016/j.jaac.2015.08.009
- **3**. Melvin GA, Dudley AL, Gordon MS, Ford S, Taffe J, Tonge BJ. What happens to depressed adolescents? a follow-up study into early adulthood. *J Affect Disord*. 2013;151(1):298-305. doi:10.1016/j.jad.2013.06.012
- 4. Leone M, Kuja-Halkola R, Leval A, et al. Association of youth depression with subsequent somatic diseases and premature death. *JAMA Psychiatry*. 2021;78(3):302-310. doi:10.1001/ jamapsychiatry.2020.3786
- **5.** Siu AL; US Preventive Services Task Force. Screening for depression in children and adolescents: US Preventive Services Task Force recommendation statement. *Ann Intern Med.* 2016; 164(5):360-366. doi:10.7326/M15-2957
- **6**. LeFevre ML; US Preventive Services Task Force. Screening for suicide risk in adolescents, adults, and older adults in primary care: US Preventive Services Task Force recommendation statement. *Ann Intern Med.* 2014;160(10):719-726. doi:10.7326/M14-0589
- 7. US Preventive Services Task Force. US Preventive Services Task Force Procedure Manual. Published 2021. Accessed August 16, 2022. https://www.uspreventiveservicestaskforce.org/uspstf/about-uspstf/methods-and-processes/procedure-manual
- 8. Viswanathan M, Wallace I, Cook Middleton J, et al. Screening for Depression, Anxiety, and Suicide Risk in Children and Adolescents: An Evidence Review for the US Preventive Services Task Force. Evidence Synthesis No. 221. Agency for Healthcare Research and Quality; 2022. AHRQ publication 22-05293-EF-1.
- 9. Forman-Hoffman V, McClure E, McKeeman J, et al. Screening for major depressive disorder in children and adolescents: a systematic review for the US Preventive Services Task Force. *Ann Intern Med.* 2016;164(5):342-349. doi:10.7326/M15-2259
- 10. O'Connor E, Gaynes B, Burda BU, Williams C, Whitlock EP. Screening for suicide risk in primary care: a systematic evidence review for the US Preventive Services Task Force. Agency for Healthcare Research and Quality; 2013.
- 11. United Nations Development Programme. Human Development Report 2019: Beyond Income, Beyond Averages, Beyond Today: Inequalities in Human Development in the 21st Century. Published 2019. Accessed August 31, 2022. https://hdr.undp.org/content/human-development-report-2019
- 12. Viswanathan M, Kennedy SM, McKeeman J, et al. Treatment of Depression in Children and Adolescents: A Systematic Review. AHRQ Comparative Effectiveness Review No. 224. Agency for Healthcare Research and Quality; 2020. AHRQ publication 20-EHC005-EF.

E12

- **13**. Sterne JAC, Savović J, Page MJ, et al. RoB 2: a revised tool for assessing risk of bias in randomised trials. *BMJ*. 2019;366:l4898. doi:10. 1136/bmj.l4898
- **14.** Sterne JA, Hernán MA, Reeves BC, et al. ROBINS-I: a tool for assessing risk of bias in non-randomised studies of interventions. *BMJ*. 2016;355:i4919. doi:10.1136/bmj.i4919
- **15.** Whiting P, Savović J, Higgins JP, et al; ROBIS Group. ROBIS: a new tool to assess risk of bias in systematic reviews was developed. *J Clin Epidemiol*. 2016;69:225-234. doi:10.1016/j.jclinepi.2015.06.005
- **16.** Whiting PF, Rutjes AW, Westwood ME, et al; QUADAS-2 Group. QUADAS-2: a revised tool for the quality assessment of diagnostic accuracy studies. *Ann Intern Med.* 2011;155(8):529-536. doi:10.7326/0003-4819-155-8-201110180-00009
- **17**. *Comprehensive Meta-analysis*. Version 3. Biostat; 2013. Accessed September 16, 2022. https://www.meta-analysis.com/
- **18**. Deeks J, Higgins J, Altman DG; Cochrane Statistical Methods Group. Analysing data and undertaking meta-analyses. In: Higgins J, Thomas J, Chandler J, Cumpston M, Li T, Page M, Welch V, eds. *Cochrane Handbook for Systematic Reviews of Interventions*. Cochrane. Published 2022. Accessed August 31, 2022. https://training.cochrane.org/handbook
- 19. US Preventive Services Task Force. US Preventive Services Task Force Procedure Manual: appendix VI: criteria for assessing internal validity of individual studies. Published 2017. Accessed August 16, 2022. https://www.uspreventiveservicestaskforce.org/uspstf/about-uspstf/methods-and-processes/procedure-manual/procedure-manual-appendix-vicriteria-assessing-internal-validity-individual-studies
- **20**. Viswanathan M, Wallace IF, Cook Middleton J, et al. Screening for anxiety in children and adolescents: evidence report and systematic review for the US Preventive Services Task Force. *JAMA*. Published October 11, 2022. doi:10.1001/jama.2022. 16303
- 21. Canals J, Bladé J, Carbajo G, Domènech-Llabería E. The Beck Depression Inventory: psychometric characteristics and usefulness in nonclinical adolescents. *Eur J Psychol Assess*. 2001;17(1):63-68. doi:10.1027//1015-5759.17. 1.63
- **22**. Christensen KS, Haugen W, Sirpal MK, Haavet OR. Diagnosis of depressed young people—criterion validity of WHO-5 and HSCL-6 in Denmark and Norway. *Fam Pract*. 2015;32(3):359-363. doi:10. 1093/fampra/cmv011
- **23.** Garrison CZ, Addy CL, Jackson KL, McKeown RE, Waller JL. The CES-D as a screen for depression and other psychiatric disorders in adolescents. *J Am Acad Child Adolesc Psychiatry*. 1991;30(4):636-641. doi:10.1097/00004583-199107000-00017
- **24.** Johnson JG, Harris ES, Spitzer RL, Williams JBW. The patient health questionnaire for adolescents: validation of an instrument for the assessment of mental disorders among adolescent primary care patients. *J Adolesc Health*. 2002;30 (3):196-204. doi:10.1016/S1054-139X(01)00333-0
- **25.** O'Connor S, Ferguson E, Carney T, House E, O'Connor RC. The development and evaluation of the Paediatric Index of Emotional Distress (PI-ED). *Soc Psychiatry Psychiatr Epidemiol.* 2016;51(1):15-26. doi:10.1007/s00127-015-1134-y

- **26.** Patton GC, Coffey C, Posterino M, Carlin JB, Wolfe R, Bowes G. A computerised screening instrument for adolescent depression: population-based validation and application to a two-phase case-control study. *Soc Psychiatry Psychiatr Epidemiol*. 1999;34(3):166-172. doi:10. 1007/s001270050129
- **27**. Roberts RE, Lewinsohn PM, Seeley JR. Screening for adolescent depression: a comparison of depression scales. *J Am Acad Child Adolesc Psychiatry*. 1991;30(1):58-66. doi:10.1097/00004583-199101000-00009
- 28. Clarke G, DeBar LL, Pearson JA, et al. Cognitive behavioral therapy in primary care for youth declining antidepressants: a randomized trial. *Pediatrics*. 2016;137(5):e20151851. doi:10.1542/peds. 2015-1851
- **29**. Clarke G, Debar L, Lynch F, et al. A randomized effectiveness trial of brief cognitive-behavioral therapy for depressed adolescents receiving antidepressant medication. *J Am Acad Child Adolesc Psychiatry*. 2005;44(9):888-898. doi:10.1016/S0890-8567(09)62194-8
- **30**. Clarke GN, Rohde P, Lewinsohn PM, Hops H, Seeley JR. Cognitive-behavioral treatment of adolescent depression: efficacy of acute group treatment and booster sessions. *J Am Acad Child Adolesc Psychiatry*. 1999;38(3):272-279. doi:10. 1097/00004583-199903000-00014
- **31.** Emslie GJ, Ventura D, Korotzer A, Tourkodimitris S. Escitalopram in the treatment of adolescent depression: a randomized placebo-controlled multisite trial. *J Am Acad Child Adolesc Psychiatry*. 2009;48(7):721-729. doi:10. 1097/CHI.0b013e3181a2b304
- **32**. Fristad MA, Vesco AT, Young AS, et al. Pilot randomized controlled trial of omega-3 and individual-family psychoeducational psychotherapy for children and adolescents with depression. *J Clin Child Adolesc Psychol*. 2019;48(suppl 1):S105-S118. doi:10.1080/15374416.2016.1233500
- **33.** Luby JL, Barch DM, Whalen D, Tillman R, Freedland KE. A randomized controlled trial of parent-child psychotherapy targeting emotion development for early childhood depression. *Am J Psychiatry*. 2018;175(11):1102-1110. doi:10.1176/appi. ajp.2018.18030321
- **34.** March J, Silva S, Petrycki S, et al; Treatment for Adolescents with Depression Study (TADS) Team. Fluoxetine, cognitive-behavioral therapy, and their combination for adolescents with depression: Treatment for Adolescents With Depression Study (TADS) randomized controlled trial. *JAMA*. 2004; 292(7):807-820. doi:10.1001/jama.292.7.807
- **35**. Mufson L, Dorta KP, Wickramaratne P, Nomura Y, Olfson M, Weissman MM. A randomized effectiveness trial of interpersonal psychotherapy for depressed adolescents. *Arch Gen Psychiatry*. 2004;61(6):577-584. doi:10.1001/archpsyc.61.6.577
- **36.** Richardson LP, Ludman E, McCauley E, et al. Collaborative care for adolescents with depression in primary care: a randomized clinical trial. *JAMA*. 2014;312(8):809-816. doi:10.1001/jama.2014.9259
- **37.** Topooco N, Berg M, Johansson S, et al. Chat- and internet-based cognitive-behavioural therapy in treatment of adolescent depression: randomised controlled trial. *BJPsych Open*. 2018;4 (4):199-207. doi:10.1192/bjo.2018.18

JAMA Published online October 11, 2022 jama.com

- **38**. Topooco N, Byléhn S, Dahlström Nysäter E, et al. Evaluating the efficacy of internet-delivered cognitive behavioral therapy blended with synchronous chat sessions to treat adolescent depression: randomized controlled trial. *J Med Internet Res.* 2019;21(11):e13393. doi:10.2196/13393
- **39**. Wagner KD, Jonas J, Findling RL, Ventura D, Saikali K. A double-blind, randomized, placebo-controlled trial of escitalopram in the treatment of pediatric depression. *J Am Acad Child Adolesc Psychiatry*. 2006;45(3):280-288. doi:10. 1097/01.chi.0000192250.38400.9e
- **40**. Lindqvist K, Mechler J, Carlbring P, et al. Affect-focused psychodynamic internet-based therapy for adolescent depression: randomized controlled trial. *J Med Internet Res.* 2020;22(3): e18047. doi:10.2196/18047
- **41.** Findling RL, Robb A, Bose A. Escitalopram in the treatment of adolescent depression: a randomized, double-blind, placebo-controlled extension trial. *J Child Adolesc Psychopharmacol*. 2013;23(7):468-480. doi:10.1089/cap.2012.0023
- **42**. Hoyniak CP, Whalen DJ, Barch D, Luby JL. Sleep problems in preschool-onset major depressive disorder: the effect of treatment with parent-child interaction therapy-emotion development. *Eur Child Adolesc Psychiatry*. 2021;30(9):1463-1474. doi:10.1007/s00787-020-01641-1
- **43**. Curry J, Rohde P, Simons A, et al; TADS Team. Predictors and moderators of acute outcome in the Treatment for Adolescents with Depression Study (TADS). *J Am Acad Child Adolesc Psychiatry*. 2006; 45(12):1427-1439. doi:10.1097/01.chi.0000240838. 78984.e2
- **44**. Emslie G, Kratochvil C, Vitiello B, et al; Columbia Suicidality Classification Group; TADS Team. Treatment for Adolescents with Depression Study (TADS): safety results. *J Am Acad Child Adolesc Psychiatry*. 2006;45(12):1440-1455. doi:10. 1097/Ol.chi.0000240840.63737.1d
- **45**. Kennard B, Silva S, Vitiello B, et al; TADS Team. Remission and residual symptoms after short-term treatment in the Treatment of Adolescents with Depression Study (TADS). *J Am Acad Child Adolesc Psychiatry*. 2006;45(12):1404-1411. doi:10.1097/01. chi.0000242228.75516.21
- **46**. Vitiello B, Rohde P, Silva S, et al; TADS Team. Functioning and quality of life in the Treatment for Adolescents with Depression Study (TADS). *J Am Acad Child Adolesc Psychiatry*. 2006;45(12):1419-1426. doi:10.1097/01.chi.0000242229.52646.6e
- **47**. McGlinchey EL, Reyes-Portillo JA, Turner JB, Mufson L. Innovations in practice: the relationship between sleep disturbances, depression, and interpersonal functioning in treatment for adolescent depression. *Child Adolesc Ment Health*. 2017;22(2):96-99. doi:10.1111/camh.12176
- **48**. Cipriani A, Zhou X, Del Giovane C, et al. Comparative efficacy and tolerability of antidepressants for major depressive disorder in children and adolescents: a network meta-analysis. *Lancet*. 2016;388(10047):881-890. doi:10.1016/S0140-6736(16)30385-3
- **49**. Thompson EA, Eggert LL. Using the suicide risk screen to identify suicidal adolescents among potential high school dropouts. *J Am Acad Child Adolesc Psychiatry*. 1999;38(12):1506-1514. doi:10. 1097/00004583-199912000-00011

- **50**. Gould MS, Marrocco FA, Kleinman M, et al. Evaluating iatrogenic risk of youth suicide screening programs: a randomized controlled trial. *JAMA*. 2005;293(13):1635-1643. doi:10.1001/jama.293.13. 1635
- **51.** Robinson J, Pan Yuen H, Martin C, et al. Does screening high school students for psychological distress, deliberate self-harm, or suicidal ideation cause distress—and is it acceptable? an Australian-based study. *Crisis*. 2011; 32(5):254-263. doi:10.1027/0227-5910/a000087
- **52.** 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): 506-514. doi:10.1016/j.jaac.2017.03.015
- **53.** 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. *Lancet Psychiatry*. 2018;5(3):203-216. doi:10.1016/S2215-0366(18) 30058-0
- **54.** Cottrell DJ, Wright-Hughes A, Collinson M, et al. A pragmatic randomised controlled trial and economic evaluation of family therapy versus treatment as usual for young people seen after second or subsequent episodes of self-harm: the Self-Harm Intervention-Family Therapy (SHIFT) trial. *Health Technol Assess*. 2018;22(12):1-222. doi: 10.3310/hta22120
- 55. Cottrell DJ, Wright-Hughes A, Eisler I, et al. Longer-term effectiveness of systemic family therapy compared with treatment as usual for young people after self-harm: an extended follow up of pragmatic randomised controlled trial. *EclinicalMedicine*. 2020;18:100246. doi:10.1016/j.eclinm.2019.100246
- **56.** 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): 122-131. doi:10.1097/00004583-201002000-00006
- **57**. 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. doi:10.1136/bmj. d682
- **58**. Griffiths H, Duffy F, Duffy L, et al. Efficacy of mentalization-based group therapy for adolescents: the results of a pilot randomised controlled trial. *BMC Psychiatry*. 2019;19(1):167. doi:10.1186/s12888-019-2158-8
- **59**. 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):662-670. doi:10.1097/CHI.0b013e3181aOacec
- **60**. Hill RM, Pettit JW. Pilot randomized controlled trial of LEAP: a selective preventive intervention to reduce adolescents' perceived burdensomeness. *J Clin Child Adolesc Psychol*. 2019;48(suppl 1):S45-S56. doi:10.1080/15374416.2016.1188705
- **61**. Hooven C, Walsh E, Pike KC, Herting JR. Promoting CARE: including parents in youth suicide prevention. *Fam Community Health*. 2012;35(3): 225-235. doi:10.1097/FCH.0b013e318250bcf9

- **62.** 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):880-893. doi:10.1037/a0016552
- **63.** King CA, Arango A, Kramer A, et al; YST Study Team. Association of the youth-nominated support team intervention for suicidal adolescents with 11-to 14-year mortality outcomes: secondary analysis of a randomized clinical trial. *JAMA Psychiatry*. 2019;76(5):492-498. doi:10.1001/jamapsychiatry. 2018.4358
- **64.** King CA, Gipson PY, Horwitz AG, Opperman KJ. Teen options for change: an intervention for young emergency patients who screen positive for suicide risk. *Psychiatr Serv*. 2015;66(1):97-100. doi:10. 1176/appi.ps.201300347
- **65**. Mehlum L, Tørmoen AJ, Ramberg M, 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):1082-1091. doi:10.1016/j.jaac. 2014.07.003
- **66.** Mehlum L, Ramberg M, Tørmoen 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. doi:10.1016/j.jaac. 2016.01.005
- **67**. Mehlum L, Ramleth RK, Tørmoen AJ, et al. Long term effectiveness of dialectical behavior therapy versus enhanced usual care for adolescents with self-harming and suicidal behavior. *J Child Psychol Psychiatry*. 2019;60(10):1112-1122. doi:10.1111/jcpp. 13077
- **68**. Haga E, Aas E, Grøholt B, Tørmoen AJ, Mehlum L. Cost-effectiveness of dialectical behaviour therapy vs. enhanced usual care in the treatment of adolescents with self-harm. *Child Adolesc Psychiatry Ment Health*. 2018;12:22. doi:10. 1186/s13034-018-0227-2
- **69**. 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):772-776. doi:10.1136/archdischild-2012-303200
- **70.** Ougrin D, Zundel T, Ng A, 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):148-153. doi: 10.1136/adc.2010.188755
- 71. 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):851-862. doi: 10.1016/j.jaac.2013.05.015
- **72.** 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):1304-1313. doi:10. 1016/j.jaac.2012.09.018
- 73. 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): 463-470. doi:10.1111/j.1440-1819.2009.01991.x
- **74.** 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):1246-1253. doi:10.1097/00004583-200111000-00003
- **75.** Psychiatry & Behavioral Health Learning Network. Beck Depression Inventory-II (BDI-II). Published 1996. Accessed April 26, 2021. https://www.hmpgloballearningnetwork.com/site/pcn/saundras-corner/scales-screenersdepression/beck-depression-inventory-ii-bdi-ii
- **76**. Rinke ML, Bundy DG, Stein REK, et al. Increasing recognition and diagnosis of adolescent depression: Project RedDE: a cluster randomized trial. *Pediatr Qual Saf*. 2019;4(5):e217. doi:10.1097/pq9.0000000000000000217
- **77**. Aboustate N, Raven M, Klau J, Jureidini J. Reanalysis of the Treatment for Adolescents with

E14

- Depression Study (TADS) under the restoring invisible and abandoned trials initiative (RIAT). *BMJ Evid Based Med.* 2019;24:A20-A21. doi:10.1136/bmjebm-2019-POD.42
- **78.** Doshi P, Dickersin K, Healy D, Vedula SS, Jefferson T. Restoring invisible and abandoned trials: a call for people to publish the findings. *BMJ*. 2013;346:f2865. doi:10.1136/bmj.f2865
- 79. US Food and Drug Administration. Suicidality in children and adolescents being treated with antidepressant medications. Published 2018. Accessed February 15, 2021. https://www.fda.gov/drugs/postmarket-drug-safety-information-patients-and-providers/suicidality-children-and-adolescents-being-treated-antidepressant-medications
- **80**. Hammad TA, Laughren T, Racoosin J. Suicidality in pediatric patients treated with antidepressant drugs. *Arch Gen Psychiatry*. 2006; 63(3):332-339. doi:10.1001/archpsyc.63.3.332

- **81.** Horowitz LM, Bridge JA, Teach SJ, et al. Ask Suicide-Screening Questions (ASQ): a brief instrument for the pediatric emergency department. *Arch Pediatr Adolesc Med.* 2012;166 (12):1170-1176. doi:10.1001/archpediatrics.2012.1276
- **82.** Aguinaldo LD, Sullivant S, Lanzillo EC, et al. Validation of the Ask Suicide-Screening Questions (ASQ) with youth in outpatient specialty and primary care clinics. *Gen Hosp Psychiatry*. 2021;68: 52-58. doi:10.1016/j.genhosppsych.2020.11.006
- **83**. The Joint Commission. Suicide prevention. Published 2021. Accessed August 16, 2022. https://www.jointcommission.org/resources/patient-safety-topics/suicide-prevention/
- **84.** National Institutes of Health. Ask Suicide-Screening Questions (ASQ) toolkit. Accessed August 16, 2022. https://www.nimh.nih. gov/research/research-conducted-at-nimh/asq-toolkit-materials

JAMA Published online October 11, 2022 jama.com