Effects of screening for psychological distress on patient outcomes in cancer: A systematic review

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Introduction

Emotional distress is common among cancer patients as a result of the diagnosis of a life-threatening disease, aggressive medical treatments, changes in lifestyle that occur, and the direct effects of the tumor [1–3]. Increasingly, attention is being paid to the psychological consequences of cancer, with recognition of not only psychiatric disorders such as major depressive disorder (MDD) or anxiety disorders, but also of subsyndromal symptoms of depression and anxiety. In addition, attention is being paid to the broader and more inclusive concept of emotional or psychological distress, as indicated by an elevated score on a one-item distress thermometer or another psychological symptom questionnaire.

A number of major cancer organizations have recommended routine screening for psychological distress in cancer care. The objective was to evaluate the effect of screening cancer patients for psychological distress by assessing the effectiveness of interventions to reduce distress among patients identified as distressed; and (2) effects of screening for distress on distress outcomes.

Methods: CINAHL, Cochrane, EMBASE, ISI, MEDLINE, PsycINFO, and SCOPUS databases were searched through April 6, 2011 with manual searches of 45 relevant journals, reference list review, citation tracking of included articles, and trial registry reviews through June 30, 2012. Articles in any language on cancer patients were included if they (1) compared treatment for patients with psychological distress to placebo or usual care in a randomized controlled trial (RCT); or (2) assessed the effect of screening on psychological distress in a RCT. Results: There were 14 eligible RCTs for treatment of distress, and 1 RCT on the effects of screening on psychological distress outcomes.

Conclusion: Treatment studies reported modest improvement in distress symptoms, but only a single eligible study was found on the effects of screening cancer patients for distress, and distress did not improve in screened patients versus those receiving usual care. Because of the lack of evidence of beneficial effects of screening cancer patients for distress, it is premature to recommend or mandate implementation of routine screening.
Well-accepted, standard definitions of medical screening define it as an intervention that involves the application of a screening tool to individuals who are not otherwise aware they are at risk, in order to detect a medical condition that can be alleviated through intervention [7,8]. Screening for MDD, for instance, involves the use of depression symptom questionnaires or small sets of questions about depression to identify patients who may have depression, but who have not sought treatment and whose depression has not already been recognized by healthcare providers. Patients identified as possible cases based on a positive screen need to be further assessed to determine if they have depression and, if appropriate, offered treatment [9].

Screening for “distress” is less well-defined since it does not seek to identify patients with a medical condition, and the meaning of a positive screen is less clear. If screening for “distress” is to be done, nonetheless, consistent with well-established definitions of screening [7,8], it would involve using scores above a pre-defined cutoff on a distress screening tool to identify patients to be offered an intervention to reduce psychological distress. Distress screening would be potentially useful if it could improve patient outcomes beyond existing standard care in which patients had access to the same services without being screened.

Three previous reviews [10–12] have sought to evaluate whether there is evidence that routine screening for psychological distress improves psychosocial outcomes among patients with cancer. The reviews have concluded that screening may improve communication between patients and health care providers and may stimulate discussions of psychosocial and mental health issues. The reviews agreed, however, that there is no conclusive evidence that screening for distress improves patient outcomes. One concern about these reviews is that they included studies that would not be considered “screening” based on any standard definition of screening. For example, some included studies used psychosocial questionnaires to inform psycho-oncology consultations that were provided to all patients. This is not screening, however, which, by definition, would involve using the questionnaires to actually determine which patients would receive the psychosocial consultations and potentially be offered psychosocial services [7–9].

In a previous systematic review, we considered the evidence on screening for MDD in cancer patients [13], but did not find evidence to support recommendations of systematic screening for depression. Compared to depression, the target of recommendations for screening for psychological distress is broader in scope, but less clearly defined in terms of targeting a specific medical condition. The objective of the present systematic review was to evaluate the evidence on screening for psychological distress in cancer. Review questions were developed based on the U.S. Preventive Services Task Force (USPSTF) [14,15] analytic framework for evaluating screening programs. The USPSTF framework recognizes the need for RCTs to directly assess links between screening programs and patient outcomes. When direct evidence from screening RCTs is not available or is of low quality, the USPSTF framework assesses key links that are necessary for screening to benefit patients, such as the availability of effective treatments [14,15].

Screening for distress per se differs from other medical screening programs in that there is not a clear, defined medical condition, such as MDD, that screening tools seek to detect. Thus, although reviews of screening usually assess screening tool accuracy compared to a gold standard [14,15], we were not able to do this. Nonetheless, an important prerequisite if screening of psychological distress is to improve patient outcomes is that distress can be reduced through intervention for patients identified as distressed. Thus, consistent with USPSTF methods, Review Question #1 was, “What are the effects of interventions to reduce distress among cancer patients who have been selected for treatment based on a minimum threshold of psychological distress, as would be done in a screening program?” If screening is to be actually recommended as policy, there should be consistent evidence from well-conducted randomized controlled trials (RCTs) [16,17] that screening benefits patients in excess of any possible harms. Thus, Review Question #2 was, “Is routine screening for psychological distress of cancer patients more effective than usual care in reducing symptoms of distress?”

Methods

Search strategy

The CINAHL, Cochrane, EMBASE, ISI, MEDLINE, PsycINFO, and SCOPUS databases were searched through April 6, 2011. A search was conducted for studies of interventions designed to reduce psychological distress among cancer patients identified as having distress (Review Question #1) and for studies that assessed outcomes of psychological distress screening interventions (Review Question #2). Search terms are reported in Appendix A. Manual searches were done on relevant systematic reviews (Appendix B), reference lists of included articles, and 45 selected journals (March 2011 to May 2012; Appendix C). We also tracked citations of included articles using Google Scholar [18] and searched the trial registries ClinicalTrials.gov [19] and the International Standard Randomized Controlled Trial Number Register [20] to attempt to identify unpublished treatment or screening RCTs.

Identification of eligible studies

Eligible articles included studies in any language on cancer patients with any type of malignancy at any disease stage that reported original data, excluding abstracts, case series, or case reports. Translators assisted reviewers to evaluate titles and abstracts and full-length articles for languages not covered by investigators, who were able to independently review material in English, Dutch, French, and Spanish. Multiple articles from the same cohort were treated as a single study. Studies with mixed populations were included only if cancer data were reported separately.

For Review Question #1, eligible articles were RCTs that compared interventions designed to reduce psychological distress to placebo, usual care, or attention controls in adult cancer patients with elevated distress. Only RCTs that limited inclusion to patients with high levels of distress, rather than all patients with cancer, were included because this is what would occur in a screening program. Indeed, patients with low levels of distress experience only negligible benefits from psychosocial interventions in cancer settings [21]. Small, underpowered studies are often of poor quality, and significant publication bias is a major problem among these studies [22–24]. A number of proposals have been made regarding setting thresholds for minimum number of patients for studies to be included in systematic reviews [23,24]. In the present review, we included trials that randomized at least 25 patients to each group [25]. Head-to-head comparisons of different interventions without a comparison to usual care or placebo were not eligible. Detailed eligibility criteria that were used for determining study eligibility are shown in Appendix D.

Eligible articles for Review Question #2 were RCTs that compared outcomes between cancer patients who underwent screening for psychological distress and those who did not. Screening was defined according to the UK National Screening Committee’s definition [7]. Thus, eligible screening trials had to include a strategy to identify patients with high levels of psychological distress based on an a priori-defined cutoff score on a measure of distress. Furthermore, in eligible studies, positive versus negative results of the screening test had to be used to make decisions about further assessment, referral, or treatment. Studies were excluded if questionnaires were used to inform and structure conversations that occurred as part of psychosocial consultations, but not to determine which patients receive services to address distress based on a score above a pre-defined cutoff. Finally, studies that involved administering multiple screening tools for multiple problems were not included, since patients in these studies could have been deemed in need of services due to reasons other than psychological distress.
(e.g., practical issues related to drug coverage by insurance, transportation and parking, or nutritional needs) [26], and determining whether the psychological distress component of screening influenced distress outcomes would not be possible.

Two investigators independently reviewed articles for eligibility. If either deemed an article potentially eligible based on title and abstract review, then a full-text review was undertaken. Disagreements after full-text review were resolved by consensus.

Evaluation of eligible studies

Two investigators independently extracted and entered data into a standardized spreadsheet (see Appendix E). Discrepancies were resolved by consensus. Risk of bias in studies included for both review questions was assessed with the Cochrane Risk of Bias tool [27] (see Appendix F), including assessment of financial conflicts of interest as has been recommended [28,29]. Risk of bias was assessed by two investigators, with discrepancies resolved by consensus.

Data presentation and synthesis

Psychological distress outcomes reported in each eligible study were classified as primary or secondary for the purposes of the review. For both review questions, when multiple measures of psychological distress were assessed as outcomes, designated primary outcomes for each study were prioritized. If there were no designated primary outcomes, the distress measure that was used to determine eligibility for the trial (Review Question #1) or as the screening tool for psychological distress (Review Question #2) was selected. If multiple instruments were used for distress selection, continuous scores on interview-based observer-rated instruments were prioritized over self-rating instruments. This is because observer-rated instruments, particularly the Hamilton Depression Rating Scale, are used most often as outcome measures in depression trials and considered the gold standard [30]. If there were no observer-rated instruments, and there was more than 1 self-rating instrument, all were reported as secondary outcomes. When outcomes were assessed at multiple time points, the assessment point that followed the end of treatment most closely was reported. Post-intervention effect sizes were reported using the Hedges’s g statistic [31], which represents a standardized difference between 2 means, as well as r², which is statistically equivalent [32,33], but presents results in terms of percent of variance in distress outcomes due to treatment. Dichotomous outcomes were not extracted since there is no agreed upon gold standard or definition for psychological distress “caseness.”

Eligible studies for each review question were evaluated to determine whether there was sufficient clinical and methodological similarity to support pooling of results. Results from trials with a high degree of clinically heterogeneity in terms of patients, interventions, or study procedures should not be synthesized meta-analytically because the effect estimate that is generated would not be expected to generalize to any given intervention [24]. For Review Question #1 (treatment), studies were heterogeneous in terms of patient samples, therapeutic interventions, outcome measures, and treatment duration. Only 1 eligible

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**Fig. 1.** PRISMA flow diagram of study selection process for review question #1.
<table>
<thead>
<tr>
<th>First author, year, country</th>
<th>Study funding source</th>
<th>Cancer type/description</th>
<th>Distress inclusion criterion</th>
<th>Treatment vs. control</th>
<th>Number of patients randomized</th>
<th>Mean age (years)</th>
<th>Males (%)</th>
<th>Treatment duration</th>
<th>Primary distress outcome: Hedges’s g (95% CI) and r²</th>
<th>Secondary distress outcome(s): Hedges’s g (95% CI) and r²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Costa, 1985, Romania [34]</td>
<td>NR</td>
<td>Mixed/mixed</td>
<td>Depression diagnosis; ZSRDS≥41; and HAMD-17 ≥16</td>
<td>Mianserin vs. placebo</td>
<td>Total: 73</td>
<td>Tx: 49 Placebo: 54</td>
<td>Tx: 0% Placebo: 0%</td>
<td>4 weeks</td>
<td>HAMD-17: 0.60 (0.13–1.07) r² = .08</td>
<td>ZSRDS: 0.64 (0.17–1.11) r² = 10 CGI-S: 0.70 (0.23–1.17) r² = .11 Δ FACT-G emotional: Δ FACT-G emotional:</td>
</tr>
<tr>
<td>Dwight-Johnson, 2005, United States [46]</td>
<td>Non-industry</td>
<td>Breast or cervical/ mixed</td>
<td>Symptoms consistent with major depression or dysthymia; or persistent depressive symptoms at baseline and 1 month later</td>
<td>Collaborative care vs. UC</td>
<td>Total: 55</td>
<td>Tx: 28 UC: 27</td>
<td>Tx: 0% UC: 0%</td>
<td>Minimum of 8 weeks</td>
<td>FACT-G emotional: 0.29 (0.05–0.54) r² = .02 SF-12 mental: 0.21 (–0.04–0.46) r² = .01</td>
<td></td>
</tr>
<tr>
<td>Ell, 2008, United States [35]</td>
<td>Non-industry</td>
<td>Mixed/mixed</td>
<td>Sadness or anhedonia more than half the days, plus PHQ-9 ≥10 and/or dysthymia</td>
<td>Collaborative care vs. enhanced UC</td>
<td>Total: 472 Tx: 242 UC: 230</td>
<td>NR</td>
<td>Tx: 16% UC: 14%</td>
<td>12 months</td>
<td>PHQ-9: 0.17 (–0.07–0.42) r² = .01</td>
<td>FACT-G emotional: 0.29 (0.05–0.54) r² = .02 SF-12 mental: 0.21 (–0.04–0.46) r² = .01</td>
</tr>
<tr>
<td>Evans, 1995, United States [36]</td>
<td>Non-industry</td>
<td>Mixed/advanced stage ii</td>
<td>CES-D ≥ 16</td>
<td>CBT vs. SSÍ vs. UC</td>
<td>Total: 78 CBT: 29 SS: 23 UC: 26</td>
<td>CBT: 54 UC: 54</td>
<td>CBT: 63% UC: 67%</td>
<td>8 weeks</td>
<td>CES-D: 0.54 (–0.02–1.10) r² = .07</td>
<td>SCL-90-R GSI: 0.59 (0.03–1.15) r² = .08</td>
</tr>
<tr>
<td>Fann, 2009, United States [37]</td>
<td>Non-industry</td>
<td>Mixed/nr</td>
<td>MDD or dysthymia diagnosis</td>
<td>Collaborative care vs. UC</td>
<td>Total: 215 Tx: 112 UC: 103</td>
<td>Tx: 72 UC: 72</td>
<td>Tx: 37% UC: 42%</td>
<td>12 months</td>
<td>SCL-20 depression: 0.47 (0.20–0.74) r² = .05</td>
<td>NA</td>
</tr>
<tr>
<td>Fisch, 2003, United States [38]</td>
<td>Non-industry</td>
<td>Mixed/advanced, incurable</td>
<td>TQSS ≥ 2</td>
<td>Fluoxetine vs. placebo</td>
<td>Total: 163 Tx: 83 Placebo: 80</td>
<td>Tx: 61 Placebo: 59</td>
<td>Tx: 55% Placebo: 45%</td>
<td>12 weeks</td>
<td>BZSDS: 0.23 (–0.21–0.66) r² = .01</td>
<td>NA</td>
</tr>
<tr>
<td>Greer, 1992, United Kingdom [39]</td>
<td>Non-industry</td>
<td>Mixed/mixed</td>
<td>HADS-D ≥ 8; HADS-A ≥ 10; or MAC helplessness ≥ 12 and MAC fighting spirit ≤ 47</td>
<td>CBT vs. UC</td>
<td>Total: 174 Tx: 85 UC: 89</td>
<td>Tx: 51 UC: 52</td>
<td>Tx: 28% UC: 14%</td>
<td>8 weeks</td>
<td>NA</td>
<td>Δ HADS-A: 0.38 (0.06–0.71) r² = .04 Δ HADS-D: 0.29 (–0.03–0.61) r² = .02 Δ MAC helplessness: 0.46 (0.13–0.78) r² = .05 Δ MAC anxious preoccupation: 0.37 (0.05–0.70) r² = .03 Δ MAC psychological: 0.33 (0.00–0.66) r² = .03 Δ PAIS psychological: 0.27 (–0.06–0.59) r² = .02 SF-36 MH quality of life: 0.32 (0.04–0.60) r² = .03</td>
</tr>
<tr>
<td>Kroenke, 2010, United States [40]</td>
<td>Non-industry</td>
<td>Mixed/mixed</td>
<td>PHQ-9 ≥ 10, plus depressed mood and/or anhedonia</td>
<td>Telecare management vs. UC</td>
<td>Total: 30953 Tx: 154 UC: 155</td>
<td>Tx: 59f UC: 59f</td>
<td>Tx: 37% f UC: 28% f</td>
<td>12 months</td>
<td>HSCL-20 depression: 0.35 (0.07–0.62) r² = .03</td>
<td>SF-36 MHI depression: 0.32 (0.04–0.60) r² = .03</td>
</tr>
<tr>
<td>Moorey, 2009, United Kingdom [41]</td>
<td>Non-industry</td>
<td>Mixed/mixed</td>
<td>HADS-A ≥ 8 or HADS-D ≥ 8</td>
<td>Nurse-delivered CBT vs. UC</td>
<td>Total: 80 Tx: 45 UC: 35</td>
<td>Tx: 65 UC: 62</td>
<td>NR</td>
<td>16 weeks</td>
<td>NR</td>
<td>NA</td>
</tr>
</tbody>
</table>
Age and sex were reported for the whole sample (N = 405), and not only the 309 participants enrolled for depression.

Treatment duration was not explicitly stated in the article, but 16 weeks was the last assessment timepoint.

Eligible participants met study criteria for depression, cancer-related pain, or both. Results are reported only for the 309 participants meeting eligibility criteria for depression.

Continued outcomes that favored the treatment group are reported as positive numbers.

Assessed using PHQ-9 and 3 additional questions from PRIME-MD.

Treatment components received varied between study participants. Effects of collaborative intervention were assessed after 8 months.

Abbreviations: BSI-GSI = Global Severity Index of the Brief Symptom Inventory; BZSDS = Brief Zung Self-Rating Depression Scale; CBT = cognitive behavior therapy; CGI-S = Clinical Global Impression Scale for Severity of Illness; CES-D = Center for Epidemiologic Studies Depression Scale; CI = confidence interval; FACT-G emotional = emotional well-being subscale of Functional Assessment of Cancer Therapy - General; HADS = Hospital Anxiety and Depression Scale total score; HADS-A = Anxiety subscale of Hospital Anxiety and Depression scale; HADS-D = Depression subscale of Hospital Anxiety and Depression Scale; HAMD-17 = 17-item Hamilton Depression Rating Scale; HAMD-21 = 21-item Hamilton Depression Rating Scale; HAS = Hamilton Anxiety Scale; HSCL-20 = 20-item Hopkins Symptom Checklist depression scale; MAC anxious preoccupation = anxious preoccupation subscale of Psychosocial Adjustment to Illness Scale; MAC helplessness = helplessness subscale of Psychosocial Adjustment to Illness Scale; MADRS = Montgomery-Asberg Depression Rating Scale; MDD = major depressive disorder; NA = not applicable; NR = not reported; PARS psychological = psychological distress subscale of Psychosocial Adjustment to Illness Scale; PHQ-9 = 9-item Patient Health Questionnaire; POMS = Profile of Mood States; PST = problem-solving therapy; PST-SO = problem-solving therapy with significant other; RSCL psychological = psychological symptoms subscale of Rotterdam Symptom Checklist; SAI = State Anxiety Inventory; SS = social support; TQSS = Two-Question Screening Survey; UC = usual care; WL = waiting list control; ZSRDS = Zung Self-Rating Depression Scale.

Mean ± SD of Mood States; HAMD-17: 3.76 (3.07–4.45) r2 = .78 PST-SO vs. WL: HAMD-17: 4.30 (3.54–5.07) r2 = .83

Total: 150

<table>
<thead>
<tr>
<th>Study</th>
<th>Industry</th>
<th>Mixed/stages</th>
<th>Distress inclusion criterion</th>
<th>Treatment</th>
<th>Number of patients</th>
<th>Average of 13 weeks</th>
<th>PST vs. WL: HAMD-17: r2</th>
<th>PST vs. WL: HAMD-21: r2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Razavi, 1996, Belgium and France [43]</td>
<td>Industry</td>
<td>Mixed/mixed</td>
<td>MDD or adjustment disorder diagnosis, plus HADS ≥ 13</td>
<td>Fluoxetine vs. placebo</td>
<td>91 Total: 91</td>
<td>Fluoxetine: 49 vs. WL: 46</td>
<td>4.45 (years)</td>
<td>4.21 (–0.12–8.48)</td>
</tr>
<tr>
<td>Strong, 2008, United Kingdom [44]</td>
<td>Non-industry</td>
<td>Mixed/mixed</td>
<td>HADS ≥ 15; MDD diagnosis; SCL-20 depression ≥ 1.75</td>
<td>Nurse intervention vs. UC</td>
<td>200 Total: 200</td>
<td>Nurse intervention: 57 vs. UC: 57</td>
<td>3.11 (−0.09–0.65)</td>
<td>r2 = .03</td>
</tr>
<tr>
<td>Van Heeringen, 1996, Belgium, [47]</td>
<td>Industry</td>
<td>Breast/stages I–II, non-metastatic</td>
<td>Depression diagnosis and HAMD-21 ≥ 15</td>
<td>Mianserin vs. placebo</td>
<td>55 Total: 55</td>
<td>Mianserin: 31 vs. placebo: 28</td>
<td>4.27 (–0.06–0.51)</td>
<td>r2 = .01</td>
</tr>
<tr>
<td>Wilkinson, 2007, United Kingdom [45]</td>
<td>Non-industry</td>
<td>Mixed/mixed</td>
<td>Depression or anxiety diagnosis</td>
<td>Aromatherapy massage vs. UC</td>
<td>288 Total: 288</td>
<td>Aromatherapy massage: 14 vs. UC: 14</td>
<td>3.87 (−0.06–0.84)</td>
<td>r2 = .01</td>
</tr>
</tbody>
</table>

Results adjusted for sex, race, years in the US, dysthymia, baseline depression severity, baseline anxiety, cancer stage, cancer type, and treatment status.

Results from social support group were not included in this review, as fewer than 25 patients were randomized to this group.

The fourth assessment visit (mean 12.3 weeks post-randomization) was closest to the end of the treatment period. However, only 33/163 patients completed the fourth visit. Outcome data presented here were assessed at the fifth visit (mean 14.9 weeks post-randomization).

Planned treatment duration was 8 weeks, but 28 patients (39%) received additional therapy sessions between 8 weeks and 4 months.

No primary outcome could be identified.

Eligible participants met study criteria for depression, cancer-related pain, or both. Results are reported only for the 309 participants meeting eligibility criteria for depression.

Age and sex were reported for the whole sample (N = 405), and not only the 309 participants enrolled for depression.

Treatment duration was not explicitly stated in the article, but 16 weeks was the last assessment timepoint.

HADS-A and HADS-D were identified in the article as primary outcomes, but insufficient information was provided to extract continuous outcome data. The authors reported that anxiety was significantly reduced in the treatment group at 16 weeks, but not depressive symptoms.

Anxiety and depression outcomes were assessed at 6 weeks.

study was identified for Review Question #2 (screening). Therefore, results were not pooled quantitatively in a meta-analysis, but were summarized in a systematic review. A review protocol was not published or registered for this systematic review. However, a written protocol was developed and followed for searching, data extraction, and data synthesis with all methods determined a priori.

**Results**

**Review question #1: effect of treatment of psychological distress**

The combined database search for Review Questions #1 (treatment) and #2 (screening) generated 4167 unique citations. As shown in Fig. 1, for Review Question #1 (treatment), 3754 were excluded after title/abstract review and 399 after full-text review leaving 14 eligible studies for review. No additional studies were identified through alternative sources, such as hand searching of journals, forward citation of included articles, and review of trial registries.

As shown in Table 1, the 14 studies of interventions to reduce psychological distress we reviewed included 12 studies of patients with mixed cancer sites [34–45], 1 study with patients with breast or cervical cancer [46], and 1 study with patients with breast cancer only [47]. Total sample size per study ranged from 55 to 472. Of the 14 studies, 7 randomized at least 64 patients per group [35,37–40,44,45], which would provide adequate (80%) power to detect a medium effect size (standardized mean difference = 0.50) [48].

Four studies were pharmacological interventions designed to treat depression, 2 with mianserin [34,47] and 2 with fluoxetine [38,43]. The other 10 studies included collaborative care interventions [35,37,40,44,46], cognitive behavior therapy [36,39,41], problem solving therapy [42], and aromatherapy massage [45]. Among the drug trials, there was 1 study [38] with at least 64 patients per group, and that study found a small effect size reduction on self-reported depressive symptoms with fluoxetine (Hedges’s g = 0.22). Three other smaller trials [34,43,47] reported somewhat larger effects for fluoxetine (Hedges’s g = 0.36) [43] and mianserin (Hedges’s g = 0.60 to 0.77) [34,47]. Among collaborative care trials, effect sizes were small to moderate for adequately powered trials (Hedges’s g = 0.17 to 0.47 [35,37,40,44]) and moderate to large for a smaller study (Hedges’s g = 0.50) [46]. The effect sizes for the replication trial of problem solving therapy trial [42], comparing problem-solving therapy to a wait-list control (Hedges’s g = 0.37) or problem-solving therapy with a significant other to the wait-list control (Hedges’s g = 4.30) were exceedingly large. The effect sizes on 2 outcome measures from aromatherapy with massage [45] were small (Hedges’s g = 0.17 to 0.22) and not statistically significant. Effect sizes for each individual study are shown in Table 1.

Risk of bias ratings are shown in Table 2, and specific explanations for all ratings are available from the authors. Among the 4 trials of antidepressants, all had unclear or high risk of bias for the majority of rating categories [34,38,43,47]. Specifically, all had unclear or high risk related to industry funding and author-industry financial ties, and all were conducted prior to the availability of clinical trial registries. Thus, selective outcome reporting was rated as unclear for all of these trials. Among non-pharmacological treatments, all were rated as high risk for blinding of patients and personnel and for blinding of outcome assessment due to the nature of the interventions and outcome assessments. Generally, quality was mixed in these studies. Not including blinding, only 1 non-pharmacological intervention trial [44] was rated as low risk of bias across all categories, including being registered with sufficiently precise outcome registration to compare to those described in the published trial report. One trial of problem-solving therapy [42] was rated as high risk of bias for Other Sources of Bias because it was due to very high sample sizes, approximately 10 times those of other non-pharmacological studies, which were reported for the primary outcome variable. Other meta-analyses have excluded this study as an extreme outlier [49–51].

**Review question #2: effect of screening for psychological distress**

For Review Question #2, 4142 of the original 4167 citations were excluded after title and abstract review and 24 after full text review, leaving 1 RCT [52] of screening for psychological distress among newly diagnosed breast cancer patients (Fig. 2). In this study, the usual care group (N = 127) received a brief psychosocial intervention as part of usual care (mean = 6.1 social work contacts versus 2.4 for usual care). As part of this study, the usual care group (N = 127) received a brief psychosocial intervention (mean 2.4 social work visits) plus telephone screening with the General Health Questionnaire, beginning 21 days post-randomization and continuing monthly for 12 months. For psychological distress, the intervention group was somewhat more likely to have a diagnosis of MDD at 12 months of follow-up (Hedges’s g = 0.36) compared to women in the control group (N = 15, 12%), although this was not statistically significant. Risk of bias in this screening RCT was generally low (Table 2).

A number of other studies (see Table 4) described by their authors or in other reviews [10–12] as related to screening were excluded from the present systematic review. Several studies were excluded because decisions about whether patients should receive further assessment, referral, or treatment were not based on a pre-specified cutoff score on a measure of distress. In those studies, a range of screening tools was often made available for clinical consultations, but a positive screen on a distress screening tool was not used to determine referral for psychosocial evaluation or treatment. Studies were also excluded because they (1) were not RCTs; (2) included multiple screening tools for many practical or logistical issues, not allowing the effect of screening for psychological distress to be evaluated separately; or (3) did not report distress symptom or diagnosis outcomes.

**Discussion**

Several clinical recommendations [4–6] have been made for screening for psychological distress to be part of standard cancer care. Guidelines and recommendations, however, vary in the degree to which they are evidence-based [53] and none of these recommendation statements have been based on a systematic review that found benefits from screening, defined according to standard definitions.

There are well-established procedures for evaluating screening programs [8,16,17]. The principal criterion is whether there is evidence from well-conducted RCTs that benefits from screening outweigh possible harms (e.g., economic costs, drug side effects). The main findings of this systematic review are that (1) treatment of distress with pharmacological or behavioral interventions can improve psychological distress in adult cancer patients with psychological distress; and that (2) only 1 RCT of distress screening, with screening defined based on standard definitions of medical screening has been conducted with adult cancer patients. In that study [52] of telephone screening for psychological distress among newly diagnosed breast cancer patients, monthly telephone screening did not improve psychological distress. The authors of that study concluded that a brief psychosocial intervention, which was provided as part of standard care, may have reduced distress and reduced the potential impact of screening. Additionally, the fact that 80% of patients in that study had at least 1 positive screen in a 12-month period suggests that screening may not have effectively identified patients with substantially elevated distress.

Several reviews on screening for distress in cancer patients have been published previously [10–12] and they each concluded that there was no evidence that distress screening improved distress outcomes among cancer patients. Two of these reviews included 7 studies [10,12], and one included 14 studies [11]. The authors of those studies were consistent in arguing that evidence for benefits of screening for distress on patient outcomes in cancer patients is inconclusive and scarce and in calling for high-quality trials to determine if distress screening would improve patient outcomes.

Two of the reviews [10,11] concluded that there is evidence that the use of distress questionnaires may improve communication about psychosocial issues between patients and oncology staff. It is important to keep in mind, however, that using questionnaires to facilitate conversations with patients, while potentially helpful, is not screening and does not inform the question of whether screening with these tools to determine who receives subsequent assessment will benefit patients. Consistent with this, a major shortcoming of previous reviews on distress screening [10–12] is that they all included studies that would not be considered trials of screening interventions in the context of any standard definition of medical screening. Indeed, with the exception of 1 study [52], all of the studies included in these reviews were excluded from the current review for a number of reasons (see Table 4 for excluded distress screening studies). Five studies [26,54–57] screened for multiple problems at the same time (i.e., fatigue, pain, perceived support, and psychological distress), which made it impossible to assess the specific effects of screening for psychological distress. One of those studies [26] screened simultaneously for multiple problems with substantially different possible care responses (e.g., psychological distress, pain, fatigue, weight change, transportation, parking, drug coverage, finances). It was not possible, however, to determine in this study how many patients screened positive for psychological distress.
versus other practical or logistical issues, such as difficulties with transportation, parking, drug coverage, or finances, none of which would be best managed through psychological intervention. Six studies [55–60] did not use a defined cutoff score to indicate a positive screen for heightened distress or to determine which patients would receive further assessment or treatment. In addition, 6 of the studies [58,60–64] were not RCTs, but were, for example, sequential cohort designs. Finally, 3 of the studies [62–64] did not assess distress as an outcome, but investigated other outcomes, such as referral rates.

Distress screening can benefit patients only to the extent that it identifies patients with significant psychological distress who are not already recognized as distressed or receiving supportive services, successfully engages those patients in treatment, and achieves positive treatment results. In many cancer care settings, however, high numbers of patients are already treated with antidepressants as an attempt to address distress, even though many of these patients do not have depression or a history of depression [65]. Furthermore, as illustrated by one study from Austria [66], the desire for psychosocial support to cope with cancer may not be correlated with distress levels, and nearly as many patients with low levels of distress may desire supportive care as patients above the cutoff criterion on a screening tool. Thus, better patient psychosocial care may be best achieved by providing more information and coordinating care pathways, rather than seeking to automate triage processes through mechanized screening and numerical algorithms.

Beyond screening for distress in cancer care settings, a number of other systematic reviews have concluded that there are no RCTs that have shown that depression screening improves depressive symptoms in cancer [13], cardiovascular disease [67], or perinatal care [68]. A 2008 meta-analysis of depression screening in primary care [69] reviewed 11 trials and found several trials where screening increased identification or treatment of depression, but none where screening improved depression outcomes. The U.S. Preventive Services Task Force has recommended depression screening in primary care [70], but specifies that screening should only occur when integrated depression care systems for evaluation and case management are available. No trials, however, have shown that patients screened and referred for such collaborative care would have better outcomes than patients who are not screened, but who could potentially access collaborative care via other pathways [9]. This was an important reason why the UK National Institute of Clinical Excellence [71] did not recommend routine depression screening in primary care.

Given the current lack of evidence for benefits of distress screening, potential costs from implementing such a program must be carefully considered. An important concern is that routine screening would either take time or consume resources that could be devoted to other patient needs. Some might assume that screening questionnaires are easily and inexpensively implemented. However, this confuses the cost of administering a questionnaire and the cost of screening. The cost of screening includes assessments, consultations,
treatment and follow-up services, which is much larger than the cost of administering a questionnaire [7,8].

Another concern is that attention and potentially limited mental health resources could be devoted only to those who screen positive for distress even though many other patients might like to discuss their psychosocial needs or might have self-referred or been referred by their clinicians. It is important that the psychological needs of cancer patients are recognized and addressed, and there are many alternatives to screening to meet this need. As long as there is no evidence that screening leads to improvements in distress, focusing on the availability and implementation of psychosocial support might better benefit cancer patients.

Without high-quality evidence from well-designed RCTs that demonstrate sufficient benefit to justify costs and potential harms from screening, recommendations for implementation of screening programs are premature. Research is needed that compares the benefits and harms of screening for psychological distress in trials in which patients in the screening group may access psychosocial resources via screening or other referral processes and patients in the non-screened group can access the same services via self- or other referral processes. Trials should clearly differentiate psychosocial needs that are best managed in the context of mental health services versus practical or logistical issues that are best addressed via other mechanisms (e.g., parking, insurance). They should also differentiate problems, such as fatigue and pain, which may or may not be related to psychological issues and for which first-line interventions are usually not psychological, from psychological distress.

**Funding/support**

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All authors have completed the Unified Competing Interest form at http://www.icmje.org/coi_disclosure.pdf and declare that Dr. Stewart is on the Depression Global Advisory Board and Cymbalta Pregnancy Registry Scientific Advisory Committee for Eli Lilly and received support from Ranbaxy Labs in 2012 to travel to Thailand to
teach on women’s mental health. No other authors have any conflict of interest disclosures for the past 3-year reporting period.

Conflict of interest

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Appendix A. Search strategy for Review Questions #1 and #2

Pubmed


Humans, clinical trial, randomized controlled trial, all adults: 19+ years

Cochrane

#1 Dep[MeSH] descriptor depressive disorder explode all trees
#2 Dep[MeSH] descriptor depression
#3 Dep[MeSH] descriptor anxiety explode all trees
#4 distress: ti,ab,kw
#5 anxiety: ti,ab,kw
#6 “quality-of-life”: ti,ab,kw
#7 (#1 OR #2 OR #3 OR #4 OR #5 OR #6)
#8 Dep[MeSH] descriptor mass screening explode all trees
#9 Dep[MeSH] descriptor psychotherapy explode all trees
#10 Dep[MeSH] descriptor treatment outcome explode all trees
#11 Dep[MeSH] descriptor antidepressive agents explode all trees
#12 Dep[MeSH] descriptor anti-anxiety agents explode all trees
#13 assess*: ti,ab,kw
#14 screen*: ti,ab,kw
#15 antidepress*: ti,ab,kw
#16 psychotherapy: ti,ab,kw
#17 psychological: ti,ab,kw
#18 (#8 OR #9 OR #10 OR #11 OR #12 OR #13 OR #14 OR #15 OR #16 OR #17)
#19 Dep[MeSH] descriptor neoplasms explode all trees
#20 cancer: ti,ab,kw
#21 tumor: ti,ab,kw
#22 tumour: ti,ab,kw
#23 oncol*: ti,ab,kw
#24 (#19 OR #20 OR #21 OR #22 OR #23)
#25 (#7 AND #18 AND #24)
#26 (randomized AND controlled AND trial): publ.type
#27 (#25 AND #26)
<table>
<thead>
<tr>
<th>First author, year, country</th>
<th>Cancer site</th>
<th>N consented/ randomized</th>
<th>Comparison</th>
<th>Distress outcomes</th>
<th>Reason(s) for exclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Berry, 2011, USA [72]</td>
<td>Mixed</td>
<td>660</td>
<td>Intervention: Patients completed a quality of life questionnaire, ESRA-C. A summary of the results, with symptoms above a predetermined threshold flagged, was provided to the clinician prior to visit. Control: Patients completed a quality of life questionnaire, the ESRA-C. No summary of the results was provided to the clinician.</td>
<td>Only number of PHQ-9 symptoms and EORTC QLQ-C30 emotional function symptoms discussed with clinician, but not distress outcomes were assessed.</td>
<td>A positive distress screen based on a defined cutoff score was not used to determine who received further assessment or treatment. Distress symptoms were not an outcome, only the discussion of symptoms.</td>
</tr>
<tr>
<td>Boyes, 2006, Australia [54]</td>
<td>Mixed</td>
<td>80</td>
<td>Intervention: Results from a computer survey completed prior to each visit were provided to the patient’s oncologist. The survey included and assessment of 12 physical symptoms associated with chemotherapy, symptoms of anxiety and depression (HADS), and perceived supportive care needs (31 items), along with computer-generated suggested strategies to manage each identified issue. Control: Results from computer survey not made available to oncologist.</td>
<td>No significant difference after 4 visits between groups for change in HADS-D scores and proportion of patients with HADS-D ≥ 11.</td>
<td>Screening of multiple problems and perceived care needs did not allow assessment of the effect of distress screening. In addition, a positive distress screen based on a defined cutoff score was not used to determine who received further assessment or treatment.</td>
</tr>
<tr>
<td>Bramsen, 2008, The Netherlands [73]</td>
<td>Mixed</td>
<td>129</td>
<td>Intervention: Patients were offered the possibility of psychosocial support by head nurse and information leaflet. Those who accepted were screened using a semi-structured interview with a checklist. Results were discussed in an interview, and patients were asked if they wanted a follow-up contact. Control: Usual care with no screening.</td>
<td>No significant difference between groups on EORTC QLQ-C30 emotional functioning subscale, IES total score or GHQ-12 total score.</td>
<td>Not a randomized controlled trial (sequential cohort design). A positive distress screen based on a defined cutoff score was not used to determine who received further assessment or treatment, which was based on whether patients requested it following an interview.</td>
</tr>
<tr>
<td>Carlson, 2010, Canada [26]</td>
<td>Lung and breast</td>
<td>1134</td>
<td>Full screening intervention: Results from DT, problem checklist, fatigue and pain thermometers, and PSSCAN, depression and anxiety sections, along with personalized feedback report placed on patient’s electronic medical record at initial visit. Triage intervention: Full screening, as described above, along with an offer to speak to a member of the study psychosocial team about any of the assessed issues. Control: DT completed, but results were not disclosed to patient or placed on electronic medical record.</td>
<td>No difference between full screening intervention, triage intervention, or usual care groups on PSSCAN depression scores 3 months post-randomization.</td>
<td>Screening of multiple problems did not allow assessment of the effect of distress screening.</td>
</tr>
<tr>
<td>Detmar, 2002, Netherlands [59]</td>
<td>Mixed</td>
<td>273</td>
<td>Intervention: Patients completed a quality of life questionnaire, the EORTC QLQ-30, at 3 successive outpatient visits with results made available to patient and physician prior to consultation. Control: Usual care.</td>
<td>No difference between groups in SF-36 Mental Health subscale after 4th visit.</td>
<td>A positive distress screen based on a defined cutoff score was not used to determine who received further assessment or treatment.</td>
</tr>
<tr>
<td>Grassi, 2011, Italy [61]</td>
<td>Mixed</td>
<td>3375</td>
<td>Intervention: Following a staff educational intervention, the DT was introduced into clinical practice, with referral to psycho-oncology services for assessment and intervention following positive screens (DT &gt; 4). Control: Physicians and nurses were able to refer patients to psycho-oncology services based on clinical judgment.</td>
<td>Only proportion of patients referred to psycho-oncology services and characteristics of referred patients were reported, but not distress outcomes.</td>
<td>Not a randomized controlled trial. In addition, distress was not an outcome. Only proportion of patients referred to psycho-oncology services was reported.</td>
</tr>
</tbody>
</table>
Hilarius, 2008, Netherlands [58]  
**Mixed 298**  
**Intervention:** Patients completed a quality of life questionnaire, the EORTC QLQ-30, at 4 outpatient visits, with summaries given to patients and nurses prior to consultation.  
**Control:** Standard consultations with physicians and nurses.  
No difference between groups in SF-36 Mental Health subscale at 4th visit.  
Not a randomized controlled trial (sequential cohort design). In addition, a positive distress screen based on a defined cutoff score was not used to determine who received further assessment or treatment.

Ito, 2011, Japan [64]  
**Mixed 998**  
**Intervention:** The pharmacist administering chemotherapy administered the DIT. Patients scoring above cutoff (≥4 distress; ≥3 impact) were recommended for consultation at psychiatry service. Those who refused were offered detailed information on self-management of mental health and were monitored. Feedback of distress results was provided on patients' medical charts.  
**Control:** Usual care with no screening  
A positive depression screen based on a defined cutoff score was not used to determine who received further assessment or treatment. In addition, screening of multiple problems did not allow assessment of the effect of depression screening.

McLachlan, 2001, Australia [55]  
**Mixed 450**  
**Intervention:** Patients at their first consultation completed a series of self-report questionnaires via touch-screen computer, including the CNQ, EORTC QLQ-C30, and BDI-SF. A summary of questionnaire results was made available to physicians prior to consultation, which were intended to be used in an individualized management plan.  
**Control:** Questionnaire responses were not made available to health care team prior to consultation.  
A positive distress screen based on a defined cutoff score was not used to determine who received further assessment or treatment.

Rosenbloom, 2007, USA [56]  
**Mixed 213**  
**Assessment, interview, and discussion intervention:** At baseline, 1 and 2 months, patients completed FLIC and FACT-G, and FACT-G scores were elaborated through an interview and discussion, the results of which were shared with the treatment nurse prior to visit.  
**Assessment intervention:** At baseline, 1 and 2 months, patients completed FLIC and FACT-G, and FACT-G scores were shared with the treatment nurse prior to visit.  
**Control:** Patients completed FLIC at baseline. Questionnaire data not shared with the treatment nurse.  
A positive distress screen based on a defined cutoff score was not used to determine who received further assessment or treatment.

Sarna, 1998, USA [74]  
**Lung 48**  
**Intervention:** Patients completed questionnaires at a number of times, including the SDS and HADS. A summary of results was given to the nurse, who identified problems and proposed interventions.  
**Control:** Patients completed questionnaires at a number of times, including the SDS and HADS. No summary of results was given to the nurse.  
SDS scores increased for the control group, but did not increase for the intervention group.  
A positive distress screen based on a defined cutoff score was not used to determine who received further assessment or treatment.

Shimizu, 2010, Japan [63]  
**Mixed 1065**  
**Intervention:** Patients completed 11-point DIT (score range 0–10), and those with a distress score ≥4 and an impact score ≥3 were referred by their oncologist for a psycho-oncology service consultation.  
**Control:** Usual care with referral to psycho-oncology services by physician of patients considered moderately or severely distressed.  
Only number of positive screens and number diagnosed and treated, but not depression outcomes, were assessed.  
Not a randomized controlled trial (sequential cohort design). In addition, outcomes included number of positive screens and number treated, but no distress outcomes were assessed.

Taenzer, 2000, Canada [60]  
**Lung 57**  
**Intervention:** At a single clinic visit, patients completed the EORTC QLQ-C30, which was provided to clinic staff prior to clinic appointment with no specific instructions for use.  
**Control:** Patients completed the EORTC QLQ-C30  
Only number of quality of life issues addressed in appointment and patients satisfaction, but no depression outcomes, were assessed.  
Not a randomized controlled trial (sequential cohort design). In addition, a positive distress screen based on a defined cutoff score was not used to determine who received further assessment or treatment and no distress outcomes were assessed.

(continued on next page)
<table>
<thead>
<tr>
<th>First author, year, country</th>
<th>Cancer site</th>
<th>N consented/ randomized</th>
<th>Comparison</th>
<th>Distress outcomes</th>
<th>Reason(s) for exclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thewes, 2009, Australia [62]</td>
<td>Mixed</td>
<td>83</td>
<td>Intervention: Patients completed the DT, and nursing staff was encouraged to assess problems and discuss psychosocial referral for patients with DT score $\geq 5$. Control: Usual care with no screening. Assessment and feedback intervention: For a 6 month study period, prior to clinic visits, patients completed the EORTC QLQ-C30 and HADS with results provided to physicians prior to visit. Attention control: For a 6 month study period, prior to clinic visits, patients completed the EORTC QLQ-C30 and HADS with no results provided to physicians. Usual care control: Patients did not complete EORTC QLQ-C30 or HADS.</td>
<td>Contrary to hypothesis, patients in the screened group reported significantly higher level of unmet needs 6 months after initial clinic contact.</td>
<td>Not a randomized controlled trial (sequential cohort design). Outcome of unmet psychosocial needs, but not distress.</td>
</tr>
<tr>
<td>Velikova, 2004, UK [56]</td>
<td>Mixed</td>
<td>286</td>
<td>Scores on FACT-Emotional Subscale were better in the intervention group than the usual care group, but not different from the attention control group.</td>
<td>Scores on FACT-Emotional Subscale were better in the intervention group than the usual care group, but not different from the attention control group.</td>
<td>A positive distress screen based on a defined cutoff score was not used to determine who received further assessment or treatment. In addition, screening of multiple problems did not allow assessment of the effect of distress screening.</td>
</tr>
</tbody>
</table>

Abbreviations: BDI-SF = Beck Depression Inventory - Short Form; CNQ = Cancer Needs Questionnaire; DIT = Distress and Impact Thermometer; DT = Distress Thermometer; EORTC QLQ-C30 = European Organization for Research and Treatment of Cancer Quality of Life Questionnaire - Core 30; ESRA-C = Electronic Self-report Assessment - Cancer; FACT = Functional Assessment of Cancer Therapy - emotional; FACT-G = Functional Assessment of Cancer Therapy - General; FLIC = Functional Living Index - Cancer; GHQ = General Health Questionnaire; HADS = Hospital Anxiety and Depression Scale; HADS-D = Depression subscale of Hospital Anxiety and Depression scale; IES = Impact of Event Scale; MDD = Major depressive disorder; PHQ = Patient Health Questionnaire; PL = Problem List; POMS = Profile of Mood States; PSI = Psychiatric Symptom Index; PSSCAN = Psychological Screen for Cancer; SF-36 = Short Form - 36 Health Survey Questionnaire.

* Number consented for non-randomized controlled trials and number randomized for randomized controlled trials.

* Physicians, rather than patients, were randomized. This number is the number of eligible patients who agreed to participate.

* Includes 2268 newly diagnosed cancer patients seen in an oncology department prior to introducing a DT and 1107 following introduction of the DT.

* The authors reported a post-hoc subgroup analysis that found significantly improved BDI-SF scores for the 44 patients in the intervention group with baseline BDI-SF scores $\geq 8$ compared to the 19 control patients with BDI SF $\geq 8$. However, patients were not randomized based on BDI-SF scores, and the relevance of these results for screening is not clear, since screening is applied to all patients, not only patients identified through screening with high scores.

* 95% of patients were female.
### Appendix B. Relevant systematic reviews


Appendix C. Coding Manual

Review Question #1: Distress treatment

Original data: The article must be an original report of a study, and not, for example, a letter, editorial, systematic review or meta-analysis, or a case series or case report study.

(Adult) cancer: The study sample must consist of cancer patients or survivors of cancer and not, for example, concern partners of cancer patients. When the sample includes cancer patients as well as other patients, data for cancer patients must be separately reported. Only studies on adult patients (≥18 years) will be included.

RCT of distress reducing intervention: The study needs to be a randomized controlled trial of treatment designed to reduce general or psychological distress as opposed to medical treatments aimed primarily at treating a physical symptom (e.g., pain, fatigue). Studies can also address treatment-specific distress, such as distress related to chemotherapy or radiation therapy. Treatments can be pharmacological, psychotherapeutic, or other. A distress treatment group has to be compared to a control group. Studies that are head-to-head comparison studies of two active treatments are not included. Only studies with placebo, standard care, or attention control are included. Studies with enhanced standard care (such as providing information to patients and/or physicians) can be included. Studies with control groups in which there is any active intervention, such as getting attention from a provider even if the attention was hypothesized to be inert, are excluded. Distress must be an outcome of the trial. Distress outcome measures can be any measure of general mental health, distress, or depression.

The following paradigm is a guide for deciding whether or not an intervention is intended to reduce distress. If a study meets at least one of the following 3 criteria, we would count it as an intervention designed to reduce psychological distress:

(1) The declared primary outcome is psychological distress (e.g., symptoms of distress, depression, anxiety, mental health function), and the intervention is not a medical treatment aimed primarily at treating the cancer (e.g., chemotherapy).

Appendix C. Journals Included in Manual Searching

Acta Psychiatr Scandinavica
American Journal of Medicine
American Journal of Psychiatry
Annals of Behavioral Medicine
Annals of Family Medicine
Annals of Internal Medicine
Archives of General Psychiatry
Archives of Internal Medicine
Australian and New Zealand Journal of Psychiatry
Biological Psychiatry
BMC Psychiatry
British Journal of Psychiatry
British Medical Journal
CA: A Cancer Journal for Clinicians
Cancer
Canadian Journal of Psychiatry
Canadian Medical Association Journal
Depression and Anxiety
European Psychiatry
General Hospital Psychiatry
Health Psychology
Herz
JAMA
Journal of Abnormal Psychology
Journal of Affective Disorders
Journal of Behavioral Medicine
Journal of Cancer Survivorship
Journal of Clinical Oncology
Journal of Clinical Psychiatry
Journal of Clinical Psychology
Journal of the National Comprehensive Cancer Network: JNCCN
Journal of General Internal Medicine
Journal of Psychosomatic Research
Journal of Supportive Oncology
Lancet
New England Journal of Medicine
New Zealand Medical Journal
Psychiatry Research
Psychological Assessment
Psychological Bulletin
Psychological Medicine
Psycho-oncology
Psychosomatic Medicine
Psychosomatics
Psychotherapy and Psychosomatics

Appendix D. Coding Manual

Review Question #1: Distress treatment
Note: If a study claims that its primary objective/outcome is to improve survival via reducing psychological distress, then count this as an intervention designed to reduce psychological distress.

(2) There are multiple outcomes declared without identification of a primary outcome, some of which are psychological and some of which are not primarily psychological (e.g., physical health or quality of life, fatigue, pain). However, the mechanism of the intervention is known to primarily target cognitions and behaviors related to mood/psychological distress or to target physiological indices of stress that are known to be related to mood/psychological distress. Examples of interventions whose mechanism is known to primarily target cognitions and behaviors related to mood/psychological distress include psychological therapies (e.g., CBT, psychodynamic therapy, behavioral therapy, expressive writing) that can be delivered via a variety of mechanisms (psychotherapy, bibliotherapy, online resources, group delivery). Coping oriented interventions would be included, as well, as coping implies a psychological component. Examples of interventions that target physiological indices of stress that are known to be related to mood/psychological distress include relaxation training, hypnosis, imagery/guided imagery, stress management, breathing training. Examples of interventions that would not meet this definition include exercise, yoga, enhanced nursing care. Note however, that all of these interventions could be included if they meet criterion 1 or 3.

(3) Criteria #1 (primary outcome) and #2 (intervention characteristics) are not met, but entry into the trial depends on meeting a threshold criteria for psychological distress. Examples of interventions in this category might include exercise, yoga, and enhanced nursing care.

Minimum level of distress: In addition, the study must include patients with a minimum level of general, psychological or emotional distress and must exclude patients scoring below that level, or studies must perform separate analyses on patients with distress scores above a cutoff level. Inclusion standards may include a self-report questionnaire or a clinical interview (structured or unstructured) for depression or anxiety disorders. Studies that do not provide separate analyses for patients above a distress cutoff, but, instead, analyze the association between distress and treatment outcome continuously are excluded. Authors will not be contacted for original data if the sample was not dichotomized in the study.

Sample size: There must be at least 25 subjects randomized to each group (distressed vs. non-distressed).

Complete distress outcome data: Outcomes have to be continuous, or a dichotomous response or remission outcome based on defined criteria must be reported.

Review Question #2: Distress screening

Original data: The article must be an original report of a study, and not, for example, a letter, editorial, systematic review or meta-analysis, or a case series or case report study.

(Adult) cancer: The study sample must consist of cancer patients or survivors of cancer and not, for example, concern partners of cancer patients. When the sample includes cancer patients as well as other patients, data for cancer patients must be separately reported. Only studies on adult patients (≥ 18 years) will be included.

RCT of screening for distress: The study needs to be a randomized controlled trial in which the intervention group patients are screened for distress with any measure or screening method and the control group is not screened. A cutoff on a distress screening tool that would be used to identify possible cases and make decisions regarding further assessment or treatment needs to be defined a priori. Studies in which questionnaire results were provided to clinicians without guidance on cutoff scores to determine positive screening status are also excluded.

Studies in which both intervention and control groups received the same psychosocial services, but service providers in the intervention group had access to results from psychosocial questionnaires that may have informed their interactions, but did not necessarily determine service allocation decisions, are excluded. Studies that administered multiple screening tools for multiple problems may be included if all of the measures have defined cutoffs for positive screens and all are screens for psychological or general distress. General or psychological distress must be an outcome of the study. Distress outcome measures can be any measure of general mental health, distress, or depression. When distress is measured, but is not an outcome variable of the study (but a predictor or mediator, etc.) studies are excluded.

Appendix E. Variables included in data extraction form

<table>
<thead>
<tr>
<th>First author</th>
<th>Year</th>
<th>Country</th>
<th>Cancer site/description</th>
<th>Distress inclusion criterion and cutoff threshold</th>
<th>Treatment condition</th>
<th>Control condition</th>
<th>N randomized, n treatment, n control</th>
<th>Mean age</th>
<th>Percentage males</th>
<th>Number and percentage lost to follow-up</th>
<th>Treatment duration</th>
<th>Distress outcomes (continuous primary and secondary outcomes):</th>
<th>Hedges’s g (95% CI) and r²</th>
<th>Study funding source</th>
</tr>
</thead>
</table>

Appendix F. Cochrane Risk of Bias Tool

**Sequence generation:** Describe the method used to generate the allocation sequence in sufficient detail to allow an assessment of whether it should produce comparable groups.

**Allocation concealment:** Describe the method used to conceal the allocation sequence in sufficient detail to determine whether intervention allocations could have been foreseen in advance of, or during, enrolment.

**Blinding of participants, personnel and outcome assessors:** *Assessments should be made for each main outcome (or class of outcomes).* Describe all measures used, if any, to blind study participants and personnel from knowledge of which intervention a participant received. Provide any information relating to whether the intended blinding was effective.

**Incomplete outcome data:** *Assessments should be made for each main outcome (or class of outcomes).* Describe the completeness of outcome data for each main outcome, including attrition and exclusions from the analysis. State whether attrition and exclusions were reported, the numbers in each intervention group (compared with total randomized participants), reasons for attrition/exclusions where reported, and any re-inclusions in analyses performed by the review authors.

**Selective outcome reporting:** State how the possibility of selective outcome reporting was examined by the review authors, and what was found.

**Pharmaceutical industry funding:** State the funding source(s) of the trial, or indicate if the trial funding source was not reported.

**Author-industry financial ties and/or employment:** State whether any trial authors disclosed financial ties and/or employment by the pharmaceutical industry, or if author-industry financial ties or affiliation were not reported.

**Other sources of bias:** State any important concerns about bias not addressed in the other domains in the tool. If particular questions/entries were pre-specified in the review’s protocol, responses should be provided for each question/entry.
References


National Collaborating Center for Mental Health. The NICE guideline on the management and treatment of depression in adults (Updated edition); 2010.

