A meta-analysis of prevalence rates and moderating factors for cancer-related post-traumatic stress disorder

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Abstract

Objective: Systematic reviews highlight a broad range of cancer-related post-traumatic stress disorder (CR-PTSD) prevalence estimates in cancer survivors. This meta-analysis was conducted to provide a prevalence estimate of significant CR-PTSD symptoms and full diagnoses to facilitate the psychological aftercare of cancer survivors.

Methods: A systematic literature search was conducted for studies using samples of cancer survivors by using validated clinical interviews and questionnaires to assess the prevalence of CR-PTSD (k = 25, n = 4189). Prevalence estimates were calculated for each assessment method using random-effects meta-analysis. Mixed-effects meta-regression and categorical analyses were used to investigate study-level moderator effects.

Results: Studies using the PTSD Checklist—Civilian Version yielded lower event rates using cut-off [73%, 95% confidence intervals (CI) = 4.5–11.7, k = 10] than symptom cluster (11.2%, 95% CI = 8.7–14.4, k = 9). Studies using the Structured Clinical Interview for Diagnostic and Statistical Manual, Fourth Edition (SCID), yielded low rates for lifetime (15.3%, 95% CI = 9.1–25, k = 5) and current CR-PTSD (5.1%, 95% CI = 2.8–8.9, k = 9). Between-study heterogeneity was substantial (I2 = 54–87%). Studies with advanced-stage samples yielded significantly higher rates with PTSD Checklist—Civilian Version cluster scoring (p = 0.05), and when assessing current CR-PTSD on the SCID (p = 0.05). The effect of mean age on current PTSD prevalence met significance on the SCID (p = 0.05). SCID lifetime prevalence rates decreased with time post-treatment (R2 = 0.56, p < 0.05).

Discussion: The cancer experience is sufficiently traumatic to induce PTSD in a minority of cancer survivors. Post-hoc analyses suggest that those who are younger, are diagnosed with more advanced disease and recently completed treatment may be at greater risk of PTSD. More research is needed to investigate vulnerability factors for PTSD in cancer survivors.

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Introduction

Systematic reviews show that long-term cancer survivorship is accompanied by co-morbid depression, anxiety and symptoms of post-traumatic stress disorder (PTSD) [1–4]. The PTSD Field Trials for the Diagnostic and Statistical Manual, Fourth Edition (DSM-IV) [5], revealed that 22% of cancer survivors present with lifetime cancer-related PTSD (CR-PTSD) [6], endorsing cancer diagnosis and treatment as a traumatic stressor. This inspired research into the prevalence and presentation of CR-PTSD in cancer populations [7].

According to DSM-IV [5], PTSD can develop when an individual experiences a traumatic event that threatens their psychological and/or physical integrity (criterion A1), and they react with helplessness or horror (criterion A2). They typically re-experience the memories, sensations and emotions from the event through sensory flashbacks and nightmares (criterion B), avoid trauma reminders and can be emotionally numb (criterion C), anxious, irritable and hyper-vigilant (criterion D). These symptoms are pathological when their duration is more than 1 month post-trauma (criterion E) and causes socio-occupational impairments (criterion F). Using the DSM-IV criteria, studies to date identified lifetime rates from 5% [7] to 35% [8]. However, the recent publication of the DSM-5 [9] challenges this position.

The new trauma criterion states ‘…a life-threatening…or debilitating medical condition is not necessarily considered a traumatic event. Medical incidents that qualify as traumatic events involve sudden, catastrophic events [e.g., waking during surgery, anaphylactic shock]’ (p. 274).

A recent article by Kangas [10] discussed how the new PTSD diagnostic criteria would affect the psychological aftercare of individuals who have completed cancer treatment—a population we will define as cancer survivors. Consequently, a PTSD diagnosis is less likely to be given to those who present with PTSD symptoms, in favour of DSM-5’s adjustment disorder (AD). AD is a diagnosis...
given to individuals who either (a) present with subsyndromal PTSD or (b) experience the complete PTSD syndrome in response to an event that is not considered traumatic [9]. Kangas [10] argued that relatively few DSM-IV studies investigated the rates of AD and PTSD in cancer samples and, thus, could not determine if PTSD was the primary disorder. This casts doubt on the appropriateness of PTSD as a diagnosis for cancer survivors but emphasises the need to correctly diagnose a cancer survivor who presents with trauma/stress-related symptoms.

It does not, however, address the needs of oncology specialists to know the proportion of patients who would likely develop PTSD or AD, or of the factors contributing to their presentation. In systematic reviews to date [1–4], efforts to achieve a prevalence estimate have been impeded owing to the substantial variability in rates, which is often attributed to extreme between-study heterogeneity in assessment methods (e.g. questionnaire versus clinical interview), assessment points (post-diagnosis or post-treatment) and sample characteristics such as severity of disease, or mean age [11].

Firstly, many studies use different methods of estimating (or diagnosing) PTSD. The most obvious example of this is the use of clinical interviews, such as the 'gold standard' Clinician-Administered PTSD Scale (CAPS) [12], or the Structured Clinical Interview for DSM-IV (SCID) [13], which yields a full DSM-IV PTSD diagnosis (criteria A–F), versus screening questionnaires such as the PTSD Checklist—Civilian Version (PCL-C) [14], which assesses symptom severity (criteria B–D). Although the PCL-C has been criticised for not establishing the fulfilment of PTSD criterion A [15], it has been administered to cancer patients and survivors in the context of experiencing cancer as a traumatic event (criterion A1). It can also assess PTSD symptoms over the past month (criterion E) and has been used frequently to assess PTSD symptom severity and provide a tentative PTSD diagnosis. Andrykowski et al. [16] validated the PCL-C against the SCID [13], and its factor structure has been corroborated in cancer survivors [17,18]. Therefore, the PCL-C is arguably a useful questionnaire to assess PTSD symptom severity and has sufficient sensitivity and specificity to screen for a probable PTSD diagnosis. Nevertheless, like other screening questionnaires, the PCL-C has been known to inflate PTSD prevalence rates owing to (a) the endorsement of PCL-C items that are confounded by disease symptoms [19]; (b) the inability to assess functional impairment (criterion F); and (c) whether cut-off scores are used or using a symptom cluster method by imitating the DSM-IV-TR PTSD diagnostic criteria [20]. The proximity of psychological assessment (by either questionnaire or structured interview) to the end of diagnosis or treatment may also contribute to prevalence. Evidence for this is equivocal; some studies have shown that PTSD persists even 20 years after the cancer experience [21], whereas others reveal no relationship between PTSD symptom severity and time since diagnosis [6,22–25] or time since treatment [26,27].

Secondly, heterogeneity may emerge from sample characteristics. Younger age at cancer diagnosis is associated with more severe PTSD symptoms [15,17,23,24,26,28–31], although some studies have not found a significant association [21,25,27]. Recent meta-analyses have demonstrated that younger samples of medical trauma survivors are associated with higher PTSD prevalence rates [32]. The disease characteristics of each study sample may also contribute. Gurevich et al. [1] argued that many studies that report no relationship between disease severity and CR-PTSD prevalence rates are actually skewed towards early disease stages [6,15,26,28–30]. However, samples that are more evenly distributed report significant effects of disease staging on prevalence rates [8,27,33,34]. In order to counter this heterogeneity, it has been argued that future studies of PTSD in cancer populations would benefit from larger, broader samples [35]. Therefore, a meta-analysis of studies reporting CR-PTSD prevalence statistics was conducted in order to find an empirical estimate of the proportion of adult cancer survivors who will present with (a) clinically significant PTSD symptoms or (b) a full diagnosis of PTSD under DSM-IV criteria. Our aims were as follows:

1. To establish a mean PTSD prevalence estimate for CR-PTSD in adult cancer survivors, with regard to the use of diagnostic interview methods and screening questionnaires. This will provide an estimate of the presentation of full-PTSD, and caseness-level PTSD symptoms indicating the need for a full assessment, respectively.

2. To establish what percentage of variance in prevalence estimates can be explained by between-study heterogeneity, with a focus on disease-stage sample characteristics, mean sample age and time since diagnosis or treatment. This will provide evidence for patient variables that may be risk factors for the development of PTSD.

**Method**

**Search strategy**

Seven online databases (MEDLINE, PsycARTICLES, PsycINFO, CINAHL Complete, CINAHL Plus, Academic Search and E-Journals) were searched systematically. Studies published from 1994 (the publication of the DSM-IV) to 11 June 2013 were included. The search terms (cancer) AND (PTSD) AND (prevalence) were used for every database. It was decided to adopt such broad terms to capture as many studies as possible that are focused on the number of CR-PTSD cases in their sample. Dissertations were included in the search, but case studies, and studies that were unavailable in English, were
excluded. Systematic reviews from 1999 onwards concerning CR-PTSD in survivors of adult cancers were examined, and any relevant references from these reviews that were not returned in online databases were abstracted. The following inclusion/exclusion criteria were compiled by the primary and secondary authors following an investigation of issues identified in the aforementioned systematic reviews and were agreed on by consensus of all the authors.

**Inclusion and exclusion criteria**

Articles were included if they met these criteria: (1) they were conducted with, and specify, a sample of cancer patients (those who are in treatment) or survivors (those who have completed treatment). Note that the search strategy does not specify any particular cancer owing to the systematic reviews arguing the scarcity of evidence for the effect of disease variables on CR-PTSD prevalence rates; (2) they provide PTSD prevalence estimates (i.e. the number of people in the sample who meet caseness); (3) the prevalence statistics are in reference to cancer diagnosis and treatment as the traumatic stressor (CR-PTSD); (4) they provide summary statistics on (a) disease stages in the sample, (b) gender, (c) mean age of sample and (d) mean time post-diagnosis or post-treatment; (5) they use both or either of the structured clinical interviews such as the CAPS [12], SCID [13], Watson’s PTSD Inventory [36] or a validated screening questionnaire that is based on DSM-IV PTSD criteria such as the PCL-C [14], or PTSD Reaction Index [37]; and (6) use cross-sectional, or longitudinal/prospective, methods. Articles were excluded if they (1) used the Impact of Events Scale as a standalone measure of probable PTSD [38]; (2) specifically used samples of adult survivors of childhood cancers; or (3) used the same sample as another included study.

**Study selection**

After the removal of duplicate entries, the full texts of the remaining records were read by the primary author. Studies that clearly met exclusion criteria (e.g. did not use cancer samples or used the Impact of Events Scale as a standalone PTSD measure) were excluded by the primary author. All authors read the remaining full-text articles and selected from the remaining studies those that they believed met inclusion criteria, and discrepancy between authors was resolved by full consensus.

**Data abstraction**

All prevalence figures, sample characteristics and study methods were abstracted onto a spreadsheet for review by the primary author and re-checked four times to ensure accurate data abstraction, and a random sample of the included studies was independently checked by a clinical psychology postgraduate (Table 1). There was full agreement. Although most of the data are easily abstracted in their presented form (e.g. mean age, gender, event rates and assessment method), some statistics were calculated manually. For example, in order to assess the moderating effect of disease-stage distribution on CR-PTSD prevalence rates, a summary statistic on the skew of the sample distribution had to be used. Given that many studies are skewed towards low disease stages, the mode (0, 1, 2, 3 or 4) was used to depict the disease stage that was most frequent in each sample. There were $k=4$ studies that were bimodal; in these cases, all bimodal distributions were characterised by I–II, II–III or III–IV, so given the nature of cancer-stage progression, the mean was used (1.5, 2.5 and 3.5, respectively). In some cases, the sample size reported in abstracts did not reflect that in the final analysis, so our abstracted $n$s reflected only those participants who completed the study. Finally, some authors reported time since diagnosis or treatment using different units (days/weeks rather than months). All studies were converted to the number of months. In all cases where it was reported in days or weeks, a month was treated as 30 days, and weeks were multiplied by seven and then divided by 30 to obtain the number of months.

**A priori statistical methods and analysis**

All statistical analyses and graphical presentations were conducted using Comprehensive Meta-analysis (CMA) [39]. Prevalence statistics were depicted using the event rate. Ninety-five percent confidence intervals (CIs) were calculated within the CMA software using the sample size ($n$) and standard error. When prospective studies were included in the analysis, the time 1 (T1) measurements (being temporally associated with study-level moderators such as time since diagnosis and/or treatment) were used to calculate the event rates. In cases where the T1 measurements were not post-treatment, the T2 prevalence estimates (and the mean time post-treatment) were used. This was the case for one study only [34]. All meta-analyses were conducted under a random-effects model owing to the substantial methodological heterogeneity identified in systematic reviews. Mixed-effects meta-regression was used to examine the association between continuous variables such as time since diagnosis or treatment end and mean age of sample, on CR-PTSD prevalence rates. Categorical analyses were conducted on studies where samples were characterised by either early (I–II) or advanced (III–IV) disease stages, and also on population studies (USA/Canada, UK/Europe and eastern countries). Finally, publication bias was assessed using funnel plots. These plots depict the spread of prevalence rates according to (a) interviews and screeners, (b) symptom cluster and cut-off scoring for screening questionnaires and (c) current versus lifetime CR-PTSD for
| Investigator | Design | Population | n | Mean age (years) | Female % | Disease stage [mode] | Cancer | Assessment point (months) | Questionnaire | Clinical interview | Cut-off | Cluster | Current | Lifetime |
|-------------|--------|------------|---|----------------|----------|---------------------|--------|--------------------------|-------------|------------------|---------|---------|---------|----------|
| Cordova et al. [26] | Cross-sectional | USA | 55 | 55.5 | 100 | I–IIIA [1] | Breast (100%) | Post-Dx | 30.5 | PCL-C | No | 5.5 | 10.9 |
| Alter et al. [6] | Cross-sectional | USA | 27 | 54 | 100 | I–II [2] | Breast (81%) | Post-Dx | 64.8 | PCL-C | No | 5 | 6 |
| Andykowski et al. [16] | Cross-sectional | USA | 82 | 56.6 | 100 | I–IIIA [1] | Breast (100%) | Post-Dx | 37.3 | PCL-C | SCID | 12 | 19 |
| Jacobsen et al. [27] | Cross-sectional | USA | 43 | 44.4 | 100 | II–IV [3] | Breast (100%) | Cut-off | 6.5 | PCL-C | No | 4.4 |
| Green et al. [7] | Cross-sectional | USA | 160 | 53.4 | 100 | I–II [1.5] | Breast (100%) | Cluster | 5.5 | PCL-C | No | 6.5 |
| Andrykowski et al. [15] | Prospective | USA | 46 | 56.4 | 100 | O–IIIA [1] | Breast (100%) | Current | 10.9 | PCL-C | No | 8.5 | 12.7 |
| Cordova et al. [17] | Cross-sectional groups (Conv versus ABMT) | USA | 142 | 56.4 (G1) | 100 | I–IIIA [1] | Breast (100%) | T1–2 | 19.5 | PCL-C | No | 14 | 15 |
| Mundy et al. [8] | Cross-sectional Groups (BMT versus Conv) | USA | 37 | 50.2 (G1) | 100 | I–IIIA [1] | Breast (100%) | T2–3 | 60 | PCL-C | No | 18 | 35 |
| Pinna et al. [25] | Cross-sectional | Israel | 50 | 52.5 | 100 | I–II [2] | Breast (100%) | Post-Dx | 19.5 | PCL-C | CAPS | 11.2 | 16.3 |
| Amiri and Ramat [59] | Cross-sectional | Israel | 39 | 50.4 | 100 | I–IIIA [1.5] | Breast (100%) | T1–2 | 60 | PCL-C | No | 2 | 117 |
| Boyer et al. [33] | Cross-sectional | USA | 133 | 56.4 | 100 | I–IIIA [1] | Breast (100%) | T1–2 | 37.2 | PTSD RI | No | 2.1 |
| Korbilz et al. [21] | Cross-sectional | USA | 153 | 55.6 | 100 | I–IIIA [2] | Breast (100%) | T1–2 | 21.6 | PCL-C | No | 4.6 |
| Lueken et al. [65] | Cross-sectional | USA | 71 | 55.6 | 100 | I–IIIA [2] | Breast (100%) | T1–2 | 6 | No | SCID | 3 |
| Palmer et al. [35] | Cross-sectional | USA | 115 | 55.6 | 100 | I–IIIA [2] | Breast (100%) | T1–2 | 12–60 | IES | SCID | 4 |
| Matsuzaka et al. [66] | Cross-sectional | Japan | 155 | 46.8 | 100 | I–IIIA [1.5] | Breast (100%) | T1–2 | 1.56 | No | SCID | 0.6 |
| Shelby et al. [19] | Cross-sectional | USA | 148 | 50.5 | 100 | I–IIIA [1.5] | Breast (100%) | T1–2 | 9.4 | PCL-C | No | 2 | 68 |
| Cordova et al. [28] | Cross-sectional | USA | 65 | 52.3 | 100 | I–IIIA [1.5] | Breast (100%) | T1–2 | 6 | PCL-C | No | 11 | 17 |
| Mehnert and Koch [67] | Longitudinal | Germany (T2) | 98 | 54.9 | 100 | I–IIIA [1.5] | Breast (100%) | T1–2 | 0.47 | PCL-C | SCID | 11.2 | 16.3 |
| Morrill et al. [24] | Cross-sectional | USA | 161 | 59 | 100 | I–IIIA [1.5] | Breast (100%) | T1–2 | 48 | No | PCL-C | No | 1.9 |
| Mehnert and Koch [68] | Cross-sectional | USA | 1083 | 61.8 | 100 | I–IIIA [1.5] | Breast (100%) | T1–2 | 46.5 | PCL-C | No | 12 | 12 |
| Sheth et al. [45] | Prospective | USA | 74 | 51 | 100 | I–IIIA [1.5] | Breast (100%) | T1–2 | 6 | PCL-C | SCID | 16.2 |
| Ganduriet et al. [89] | Case-control | France | 144 | 53 | 100 | I–IIIA [2] | Breast (100%) | T1–2 | 21 | No | Watson's PTSD Inventory | 49 | 2.8 |
| Poskum et al. [74] | Longitudinal | USA | 53 | 50 | 100 | I–IIIA [2] | Gynaecological | T1–2 | 6.5 | PCL-C | No | 22.4 |
| Goncalves et al. [42] | Longitudinal | Portugal (T1) | 161 | 61 | 100 | I–IIIA [2] | Ovarian | T1–2 | 2.8 | No | SCID | 23.4 |
| Mistakidou et al. [41] | Cross-sectional | Greece | 989 | 64.4 | 49.2 | III–IVIA [1.5] | Breast (18%) | T1–2 | 0 | No | SCID | 11.4 |

Time points—T1, T2 and T3. Group—G1, G2 and G3.
ABMT, autologous bone marrow transplant; Conv, conventional treatment.
clinical interviews. In order to calculate the variance contribution of statistically significant study-level moderators on PTSD event rates, we calculated the $R^2$ statistic using the following formula: 

$$R^2 = 1 - \left( \frac{T^2_{\text{unexplained}}}{T^2_{\text{total}}} \right).$$

**Results**

**Literature search and study characteristics**

We identified 257 articles. After duplicates were removed, 109 articles remained. A combined total of 138 studies were recovered from online databases and systematic reviews. Figure 1 depicts the PRISMA search strategy [40]. Table 1 presents the abstracted prevalence data and study/sampling characteristics for the final dataset ($k=25$). The aggregated sample was $n=4189$, of which 88% were female, and the study of Mystakidou et al. [41] was the only study to represent both genders. In terms of methodology, $k=19$ studies were cross-sectional, and $k=6$ were longitudinal/prospective. The majority of studies were conducted in the USA ($k=18$), with one in Japan, five in Europe and the UK, and one in Israel. Although there was a range of cancers included, 21 studies used exclusively female breast cancer samples, with one study using ovarian cancer [42], one gynaecological cancer [34] and two inclusive of breast cancer, but with the addition of others [6,41]. Owing to the low $n$ for cancers other than breast, an analysis of the moderating effects of cancer type could not be conducted. Therefore, a separate analysis was conducted on breast cancer alone, as this is the most frequently studied cancer population. Also, the moderating effects of gender were not analysed owing to the predominance of women over men in the final dataset.

**CR-PTSD prevalence rates**

The results of the meta-analysis are available in Table 2. The table presents prevalence statistics for the whole pool of cancer studies and for breast cancer alone, grouped by
assessments by the PCL-C cluster method and the SCID yielded remarkably similar lifetime prevalence estimates (13.8%, 95% CI = 9.5–19.6, k = 11). No comparison could be performed between interview methods, or between screening tools, because only the PCL-C and the SCID were used more than once. Although the point estimates for each assessment method are comparable, each is characterised by considerable between-study heterogeneity across cancer samples (Table 2; $I^2 = 54.4–81.5\%$).

Breast cancer: Studies that used the PCL-C yielded lower event rates using the cut-off method (6.4%, 95% CI = 4.2–9.7, k = 9) compared with the symptom cluster method (11.2%, 95% CI = 8.7–14.4, k = 9). These figures are similar to those found across cancers, although the cut-off event rate for breast cancer alone is 1% less. The rates for the symptom cluster method here are the same for breast cancer as for all cancers. Studies using the SCID alone yielded lower but nevertheless comparable event rates for current CR-PTSD (4.1%, 95% CI = 2.8–8.5, k = 7). Much higher rates were found for lifetime CR-PTSD (14.2%, 95% CI = 7.7–24.9, k = 4). Studies that used a clinical interview method yielded a combined event rate of 5.8% (95% CI = 3.3–10, k = 10) for current CR-PTSD and 11.5% (95% CI = 6.3–20.1, k = 6) for lifetime CR-PTSD. Studies using the cut-off scoring method used the PCL-C exclusively (see aforementioned results). All other screening tools used a symptom cluster method and yielded remarkable similar prevalence estimates to the PCL-C cluster method (12.1%, 95% CI = 9.3–15.7, k = 10). Similar between-study heterogeneity was observed for breast cancer samples ($I^2 = 54.4–81.5\%$).

### Sources of heterogeneity

Mean sample age, mean time since diagnosis and mean time since end of treatment were entered into a meta-regression for their variance contribution to CR-PTSD prevalence rates. Early or advanced modal disease staging and population of origin (USA, UK and Europe, Japan and Israel) were analysed categorically to assess group differences in prevalence.

### Table 2. PTSD event rates (a) across cancers and (b) for breast cancer samples, according to method of assessment

| Cancer-related PTSD measure | k | Event rate | 95% CI | $p$ | Q | d.f. | $i^2$ | $\tau^2$ | $\tau^2$ | SE | Variance |
|-----------------------------|---|-----------|-------|-----|---|-----|------|--------|-------|-----|---------|
| **Breast cancer (Table 2; between-study heterogeneity across cancer samples are comparable, each is characterised by considerable between-study heterogeneity across cancer samples)** | | | | | | | | | | | |
| **By assessment method** | | | | | | | | | | | |
| Interview | Current | 10 | 5.8 | 3.3 | 10 | <0.01 | 33.4 | 9 | 73.05 | <0.01 | 0.61 | 0.78 | 0.42 | 0.18 |
| | Lifetime | 6 | 11.5 | 6.3 | 20.1 | <0.01 | 27.01 | 5 | 81.5 | <0.01 | 0.53 | 0.73 | 0.44 | 0.19 |
| Questionnaire | Cut-off | 9 | 6.4 | 4.2 | 7.9 | <0.001 | 17.56 | 8 | 54.44 | <0.01 | 0.24 | 0.49 | 0.23 | 0.06 |
| | Cluster | 10 | 12.1 | 9.3 | 15.7 | <0.01 | 28.14 | 9 | 68.02 | <0.01 | 0.14 | 0.37 | 0.12 | 0.01 |
| **By specific tool** | | | | | | | | | | | |
| SCID | Current | 7 | 4.1 | 2 | 8.5 | <0.001 | 21.29 | 6 | 72.59 | <0.001 | 0.7 | 0.83 | 0.61 | 0.37 |
| | Lifetime | 4 | 14.2 | 7.7 | 24.9 | <0.001 | 15.35 | 3 | 80.7 | <0.001 | 0.4 | 0.63 | 0.42 | 0.17 |
| PCL-C | Cut-off | 9 | 6.4 | 4.2 | 9.7 | <0.001 | 17.56 | 8 | 54.44 | <0.03 | 0.24 | 0.49 | 0.23 | 0.06 |
| | Cluster | 9 | 11.2 | 8.7 | 14.4 | <0.001 | 18.8 | 8 | 57.45 | <0.02 | 0.1 | 0.31 | 0.1 | 0.01 |
| **Across cancers** | | | | | | | | | | | |
| **By assessment method** | | | | | | | | | | | |
| Interview | Current | 12 | 6.4 | 4.1 | 9.9 | <0.01 | 42.8 | 11 | 74.29 | <0.01 | 0.41 | 0.64 | 0.34 | 0.11 |
| | Lifetime | 7 | 12.6 | 7.4 | 20.7 | <0.01 | 28.73 | 6 | 79.2 | <0.01 | 0.49 | 0.7 | 0.38 | 0.15 |
| Questionnaire | Cut-off | 10 | 7.3 | 4.5 | 11.3 | <0.001 | 31.4 | 9 | 71.33 | <0.01 | 0.47 | 0.68 | 0.33 | 0.11 |
| | Cluster | 11 | 13.8 | 9.5 | 19.6 | <0.01 | 75.89 | 10 | 86.82 | <0.01 | 0.41 | 0.64 | 0.29 | 0.08 |
| **By specific tool** | | | | | | | | | | | |
| SCID | Current | 9 | 5.1 | 2.8 | 8.9 | <0.001 | 35.46 | 8 | 77.43 | <0.001 | 0.53 | 0.72 | 0.49 | 0.24 |
| | Lifetime | 5 | 15.3 | 9.1 | 24.7 | <0.001 | 16.64 | 4 | 75.96 | <0.001 | 0.34 | 0.59 | 0.33 | 0.11 |
| PCL-C | Cut-off | 10 | 7.3 | 4.5 | 11.7 | <0.001 | 31.4 | 9 | 71.33 | <0.01 | 0.47 | 0.68 | 0.33 | 0.11 |
| | Cluster | 9 | 11.2 | 8.7 | 14.4 | <0.001 | 18.8 | 8 | 57.45 | <0.09 | 0.31 | 0.01 | 0.01 | 0.01 |

**Notes:** The rates for the symptom cluster method here are similar to those found across cancers, although the cut-off event rate for breast cancer alone is 1% less. The rates for the symptom cluster method here are the same for breast cancer as for all cancers. Studies using the SCID alone yielded lower but nevertheless comparable event rates for current CR-PTSD (4.1%, 95% CI = 2.8–8.5, k = 7). Much higher rates were found for lifetime CR-PTSD (14.2%, 95% CI = 7.7–24.9, k = 4). Studies that used a clinical interview method yielded a combined event rate of 5.8% (95% CI = 3.3–10, k = 10) for current CR-PTSD and 11.5% (95% CI = 6.3–20.1, k = 6) for lifetime CR-PTSD. Studies using the cut-off scoring method used the PCL-C exclusively (see aforementioned results). All other screening tools used a symptom cluster method and yielded remarkable similar prevalence estimates to the PCL-C cluster method (12.1%, 95% CI = 9.3–15.7, k = 10). Similar between-study heterogeneity was observed for breast cancer samples ($I^2 = 54.4–81.5\%$).
Time post-diagnosis was not significant for any analyses. Time post-treatment was not significant when using the SCID, PCL-C or the questionnaire method but was significant when including other clinical interviews to assess lifetime CR-PTSD \( Q_{(1)} = 3.84, \ T^2_{\text{unexplained}} = 0.22 \text{ k = 3, } p = 0.05 \), with CR-PTSD event rates decreasing when time since treatment increases, and a variance contribution of 56\% \( (R^2 = 0.56, p < 0.05) \). The contribution of mean sample age to prevalence was not significant for breast cancer samples, or when using the cut-off and cluster screening methods and the clinical interview method, or when using the SCID to assess lifetime CR-PTSD in all cancers. However, it was significant when using the SCID to assess current CR-PTSD \( Q_{(1)} = 0.43, p = 0.05 \).

Disease stage was significant when using the SCID to assess current CR-PTSD \( Q(1) = 10.23, p = 0.05 \). Studies characterised by early stages (I–II) yielded markedly lower prevalence rates (4.2\%, 95\% CI = 2.1–8.1, \( k = 8, n = 737 \)) than those with advanced-stage (III–IV) disease (11.4\%, 95\% CI = 9.6–13.5, \( k = 11, n = 989 \)). This was also true for the interview method \( Q(1) = 6.07, p = 0.01 \), with the only difference in effect coming from the additional interview methods other than the SCID in the early-stage group (5.7\%, 95\% CI = 3.4–9.6, \( k = 11, n = 970 \)). Finally, the use of the questionnaire method with cluster scoring also yielded a significant difference \( Q(1) = 3.71, p = 0.05 \) with early stages yielding significantly lower prevalence rates (11.6, 95\% CI = 8.8–13.5, \( k = 9, n = 1894 \)) than advanced-stage samples (31.3 95\% CI = 11.8–61, \( k = 2, n = 112 \)). This moderator was not significant for pure breast cancer samples.

Population was not a significant moderator when using the questionnaire method or on the PCL-C, or the SCID, alone, and in either the full cancer sample or breast cancer samples. But significant differences were found when using the clinical interview method to diagnose current CR-PTSD \( Q(3) = 9.2, p = 0.03 \), with Israel having significantly higher rates (18\%, 95\% CI = 8.8–33.2, \( k = 1 \)) than the USA (5.6\%, 95\% CI = 2.6–11.6, \( k = 8 \)), Europe (6\%, 95\% CI = 2.4–14, \( k = 3 \)) or Japan (3.9\%, 95\% CI = 1.8–8.4, \( k = 1 \)). Population was also significant for lifetime CR-PTSD \( Q(2) = 11.28, p = 0.03 \), with the USA (15.4\%, 95\% CI = 8.4–26.7, \( k = 5 \)) and Japan (14.8\%, 95\% CI = 10–21.3, \( k = 1 \)) having higher rates than Europe (2.8\%, 95\% CI = 1.1–7.2, \( k = 1 \)). Population was significant in exclusively breast cancer samples when diagnosing lifetime CR-PTSD with the interview method \( Q(2) = 10.83, p < 0.01 \), with the UK and Europe (2.8\%, 95\% CI = 1.1–7.2) having a significantly lower prevalence than the USA (14.2\%, 95\% CI = 6.8–27.5, \( k = 4 \)) and Japan (14.8, 95\% CI = 10–21.3, \( k = 1 \)). Similar trends in breast cancer were found when diagnosing current CR-PTSD \( Q(3) = 11.43, p < 0.01 \), with Israel yielding higher rates (18\%, 95\% CI = 8.8–33.2, \( k = 1 \)) than Japan (3.9\%, 95\% CI = 1.8–8.4, \( k = 1 \)), UK and Europe (4.1\%, 95\% CI = 2.2–7.6, \( k = 2 \)) and USA (5.7\%, 95\% CI = 2.5–12.6, \( k = 6 \)).

**Discussion**

This is thought to be the first meta-analysis to investigate the prevalence of CR-PTSD in cancer survivors, and the contribution of between-study heterogeneity to the fluctuation of these rates. Our data for all cancers indicated that studies using clinical interview methods yield a mean prevalence of 6.4\% for current CR-PTSD and 12.6\% for lifetime CR-PTSD. Screening methods that used cluster scoring indicated a prevalence of 13.8\% for clinically significant CR-PTSD symptoms. The PCL-C yielded lower event rates of clinically significant CR-PTSD symptoms using the cut-off (7.3\%) compared with symptom cluster (11.2\%), whereas the SCID yielded predictably higher lifetime diagnoses (15.3\%) than current diagnoses (5.1\%) for CR-PTSD.

These rates are similar to those found by Alter et al. [6] in the DSM-IV Field Trials (4\%), Green et al. [7] and Palmer et al. [35], which identify a current CR-PTSD prevalence of 2.5–5\%. Epidemiological surveys reveal the lifetime conditional risk (CR) of PTSD for women in the US general population to be between 10.4\% (95\% CI = 8.8–11.7\%) [43] and 13\% (95\% CI = 9.9–16.1\%) [44]. Nevertheless, the lifetime CR-PTSD rate for our 100\% female US sample was 15.4\% (95\% CI = 8.4–26.7\%), which is higher than the CR for the USA, but within the CIs for Breslau’s survey [44]. If 5–12\% of cancer survivors develop CR-PTSD, a trauma framework may not represent the distress experienced by most patients—as argued by Green et al. [7], Palmer et al. [35] and now the DSM-5 task force [9]. This supports Kangas’s criticism that the lack of differential diagnoses in semi-structured clinical interviews challenges the validity of PTSD diagnosis [10]. It does not, however, take into account that the literature fails to record the presence/absence of discrete catastrophic events within the course of the disease that would make the cancer experience traumatic.

We recommend that clinicians consider the full range of presentations, be aware of the nature of cancer as a stressor and assess for discrete events that occurred during the treatment to warrant a PTSD diagnosis and, in the absence of these, consider a diagnosis of AD. If cancer survivors meet AD criteria, this does not discount the possibility of PTSD development. The literature shows that cancer survivors present with subsyndromal symptoms [7,45,46], which are also associated with considerable functional impairments [26,45]. Although this too may lead clinicians to initially question the validity of a CR-PTSD diagnosis, it does not discount the possibility that these subsyndromal symptoms may be a predictor of future PTSD. DSM-5 course specifiers state that PTSD may present with delayed expression, where only
This relationship concentrates on age at diagnosis and the steady accumulation of subsyndromal symptoms from the onset of the stressor [47,48]. These are features of cancer, and subsyndromal symptoms are also present at the beginning of cancer treatment, fluctuate and persist throughout the course of the disease and may, too, reflect a delayed PTSD presentation [15]. Although rightly considered an adjustment response during treatment, this may develop into CR-PTSD if those factors prevent the cognitive processing of the illness.

In our moderator analysis, screening questionnaires yielded higher rates of CR-PTSD symptoms than did the use of clinical interviews for current CR-PTSD. The reasons for this may be that whereas interviews diagnose disorder, questionnaires detect clinically significant symptoms, are not clinician administered and do not screen for disorders that better explain the symptoms [20]. However, the PCL-C has demonstrated its reliable sensitivity and specificity in correctly identifying clinical cases [11], but there is still debate on where to place the cut-off score [20]. The Davidson Trauma Scale [49] can be used as an alternative to screen for CR-PTSD. It has a cut-off score that moves depending on the prevalence of PTSD in the specific population. As this meta-analysis revealed a current CR-PTSD prevalence of 6.4%, the Davidson cut-off for the cancer survivor population would be 47.

When lifetime CR-PTSD was assessed using the SCID, prevalence decreased with time post-treatment. This finding should be interpreted with caution. Firstly, this trend was only observed for lifetime CR-PTSD, and secondly, this factor is a proxy variable for time post-trauma. As there is no agreement as to whether the diagnosis or the treatments are the definitive traumas, investigators adopted one of the two indices; therefore, these moderators could not be combined, resulting in low statistical power. However, this finding reflects the epidemiologic trend that non-cancer PTSD sufferers do recover naturally over time, although a minority of them remain symptomatic after many years [43]. This has implications for the survivors, as potential cases may not be identified during routine follow-up, placing them at risk of PTSD months to years after their treatment.

Younger-aged samples were associated with higher current CR-PTSD event rates on the SCID. This trend was just significant, so these results too must be interpreted cautiously. The majority of research into this relationship concentrates on age at diagnosis—our study-level moderator was the mean age of the sample. Several of the samples were many months, or sometimes years, post-diagnosis, and this likely reduced the sensitivity of our analysis—but as it addressed the impact of younger age on the prevalence rates of lifetime CR-PTSD (not on symptom severity)—it provides a singular contribution.

The final aim was to synthesise disease-stage data to establish whether they are a risk factor for PTSD following comments that skewed samples characterised the research [1]. Our post-hoc analyses suggested that advanced disease is related to an increase in CR-PTSD event rates on PCL-C and on the SCID. This is commensurate with studies that show a positive relationship between disease severity and CR-PTSD [8,27,33,34] and may provide some preliminary support for this conclusion, but we advise cautious interpretation. Although the differences in event rates were significant, additional factors may have contributed. The advanced disease sample in one study [41] was from a different culture and set in palliative care, which introduces environmental factors into the development of CR-PTSD that are uncharacteristic of early-stage cancer samples. The degrees of freedom for the early-stage group were larger than those for the advanced-stage group and reflect the abundance of earlier stage cancers in the majority of studies.

Publication bias

The funnel plots revealed substantial biases. However, this is not the bias one would expect from the file-drawer problem. Publication bias is identified by a skew in the distribution towards higher effect sizes, at the bottom of the funnel plot. The presence of this type of bias is not visually depicted in our plots.

Strengths and limitations

There are limitations that constrain our conclusions. The findings from the moderator analysis are tentative because several findings come from retaining $k = 1$ comparisons. This is problematic as the $Q$ test has low power to detect heterogeneity when $k$ is low [50]. Nevertheless, this highlights the need for concise, standardised and transparent reporting to facilitate future meta-analytic studies. Also, only study-level moderators were included. Many studies did not assess psychiatric history or additional life stress at the time of assessment, so intrapersonal vulnerability factors were unaccounted for. The substantial variability in reporting styles, and limitations of using CMA, may have introduced variance into analyses that is not attributable to the moderators of interest. Studies that have used questionnaires may over-inflate rates of CR-PTSD owing to symptom endorsement being confounded by artefacts of cancer drugs, and medical conditions [5], and realistic fears of cancer recurrence being endorsed as the acceptance of a foreshortened future. Nevertheless, the sensitivity and specificity of the PCL-C against the SCID for CR-PTSD are sufficient to screen for those survivors who may be suffering from an AD or CR-PTSD. Out of the studies that have used interviews, few have assessed the rates of co-morbid...
disorders, meaning CR-PTSD might not be the primary disorder. Also, all of the studies included in this meta-analysis used DSM-IV criteria, not the new DSM-5. Early epidemiological studies documented the inflation of PTSD prevalence owing to the revised DSM-IV criteria, so PTSD prevalence may decrease owing to the new DSM-5 PTSD criterion A. The strengths of this investigation are that it included 25 studies and a substantial patient pool across several populations. This affords our analysis generalisability that the individual studies could not achieve [35]. Precision was enhanced by including data for symptoms anchored to the experience of adult cancer. This does not account for differential diagnoses but does account for stress-related symptoms due to an extreme stressor. In this case, the findings may reflect the proportion of survivors who meet criteria for AD and PTSD. This is more clinically useful than focusing on PTSD alone.

Future research

If the cancer experience is not generally traumatic, then the fact that CR-PTSD does emerge in a significant minority of survivors suggests that other factors affect adjustment and the cognitive processing of the illness. It has been argued that ‘less traumatic’ stressors can be expected to trigger PTSD in vulnerable individuals who have limited resources to manage their response to stress [51] and that the insufficiency of some events to meet PTSD criterion A may reflect failure to consider the role of personal stress reactivity [52,53], and the effect of allostatic load [53]. As such, there is less information on variables that influence this adjustment. Variables such as social support and/or constraints on talking about the illness [21,28,30,54–58], emotion regulation strategies such as dissociation [22], emotional suppression and avoidance [59,60], and rumination [61] have been identified as predictors of PTSD symptoms in cancer populations. Most recently, emotional processing styles have been implicated in post-natal depression [62] and PTSD [63,64]. Given the protracted nature of cancer, it is possible that social variables and their relationship to beliefs about experiencing and expressing of emotions may influence cognitive processing over the course of disease and predict the emergence of CR-PTSD. This necessitates further investigation on the role of these variables in multiple oncology services both nationally and internationally. But firstly, in order to assess the risk of CR-PTSD in this population, it may be necessary to report, or make readily available, the raw data on CR-PTSD event rates by (a) disease stage, (b) cancer type, (c) treatment regimens, (d) any traumatic events during the course of the disease, (e) age at diagnosis and (f) the mean time of assessment in relation to time post-diagnosis and time post-treatment.

Conclusions

Prevalence rates from questionnaires reveal that a minority of cancer survivors present with clinically significant symptoms due to cancer. In these cases, we recommend that oncology specialists recognise the possibility of CR-PTSD and refer their patient to mental health services. Prevalence rates from clinical interviews show that a minority of cancer survivors meet (or have previously met) full DSM-IV criteria for CR-PTSD after the conclusion of treatment and that those who are younger, diagnosed with more advanced disease or recently completed treatment may be at greater risk of CR-PTSD. Methodological heterogeneity prohibits robust conclusions about the expected prevalence of CR-PTSD as a primary disorder but does provide some clinical justification for the diagnosis of a CR-PTSD in a minority of cases. Given the release of the DSM-5, we recommend caution in diagnosing CR-PTSD but advise further investigation into whether traumatic occurrences were experienced during the course of the disease.

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Conflict of interest

The authors have declared no conflicts of interest.

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