

REVIEW

Limited evidence exists on the effectiveness of education and training interventions on trial recruitment; a systematic review

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Abstract

Objective: The objective of this study was to examine the effectiveness of education and training interventions on recruitment to randomized and non-randomized trials.

Study Design and Setting: A systematic review of the effectiveness of education and training interventions for recruiters to trials. The review included randomized and non-randomized controlled trials of any type of education and training intervention for recruiters to trials, within any health care field. The primary outcome was recruitment rates, and secondary outcomes were quality of informed consent, recruiter self-confidence, understanding/knowledge of trial information, numbers of potential trial participants approached, satisfaction with training, and retention rates.

Results: Of the 19 records reviewed at full-text level, six met the inclusion criteria for our review. Owing to heterogeneity of outcomes and methods between the included studies, meta-analysis was not possible for the primary outcome. Of the three studies that reported recruitment rates, one favored the education and training intervention for increased recruitment; the remaining two found no differences between the groups. Of the reported secondary outcomes, quality of informed consent was improved, but no differences between groups in understanding/knowledge of trial information were found.

Conclusion: There is limited evidence of effectiveness on the impact of education and training interventions on trial recruitment. Further work on developing a substantial evidence base around the effectiveness of education and training interventions for recruiters to trials is required. © 2019 Elsevier Inc. All rights reserved.

Keywords: Trial recruitment; Educational intervention; Training intervention; Systematic review; Effectiveness review; Trial methodology

1. Background**1.1. Introduction**

Randomized control trials are considered the gold standard for testing the effectiveness of interventions. However, trialists face many challenges in trial processes. One major challenge is recruitment, with reports suggesting that fewer than half of trials reach their original recruitment targets or require an extension to the trial to do so [1].

In a survey of authors of published primary care trials, less than one-third reported that they recruited to target within the original timescale [2]. In addition, McDonald et al. [3] explored levels of recruitment in a cohort of 114 trials in the United Kingdom (from 1994 to 2002) and found that recruitment problems were identified in the early stages of 63% of the trials, only 31% achieved their original recruitment target and 53% were given an extension. More recently, Sully et al. [4] concluded that although recruitment has improved since McDonald's study, only half (55%) of the trials in their study recruited the original target sample size and 45% were extended.

Poor recruitment has many negative consequences, for instance, trial extensions are often required to reach

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What is new?

Key findings

- There is limited evidence overall, on the effectiveness of education and training interventions on recruitment to trials, and the evidence that exists is conflicting.
- Education and training interventions, however, have demonstrated improved quality of informed consent.

What this adds to what was known?

- Further work on developing a substantial evidence base around the effectiveness of education and training interventions for recruiters to trials is required.
- Outcome measures and methods of measuring outcomes across studies are varied, making it difficult to synthesize, and compare and contrast study's findings.

What is the implication and what should change now?

- A standardized set of outcome measures is required for use in future evaluations of trial recruitment education and training interventions.

recruitment targets, these are costly [5,6]. If recruitment targets are not met, the research question often remains unanswered, wasting money, much research effort, and participants' time. Strategies are often implemented to improve recruitment rates such as offering incentives to potential participants or tailored recruitment materials. However, as highlighted by Treweek et al. [1], there is limited high-quality evidence on whether or not particular recruitment strategies are effective. They found high-certainty evidence for three recruitment strategy comparisons (of 72) and concluded that further evaluation and replication of evaluations are required to strengthen the evidence base.

1.2. Rationale

This review is part of a wider study (Training Recruiters—An educational Intervention [TRAIN]), which aims to develop and evaluate an education and training intervention for recruiters to trials. A previous review of training programs for recruiters to trials, by Townsend et al. [7], reviewed all study types (e.g., qualitative, pre-test/post-test, randomized and non-randomized), assessed their quality using the Effective Public Health Practice Project tool, and summarized the results narratively. They found that some training interventions increased recruiter

self-confidence when communicating trial information, but found little evidence that interventions increased recruitment rates, informed consent, patient understanding, and satisfaction. They concluded that further development of training interventions for trial recruiters, with a focus on improving recruitment and informed consent, is required.

The current review was carried out to provide contemporary, up-to-date evidence on the effectiveness of education and training interventions for recruiters to trials, so as to inform the design of the TRAIN. Although some studies included in the review of Townsend et al. [7] were likely eligible for inclusion in our review, our review, in addition to searching for new trials, includes randomized and non-randomized control trials only which evaluated education and training interventions within the context of a planned or an ongoing trial (“host trial”) and reported outcomes of effectiveness.

1.3. Aims and objectives

The objective of this study was to determine the effectiveness of education and training interventions on recruitment to randomized and non-randomized control trials (here after referred to as trials). Our primary objective is to explore whether or not training and educational interventions for recruitment to trials positively affects recruitment rates.

2. Methods

2.1. Protocol and registration

The review protocol is registered on PROSPERO (ID = CRD42018108019) and can be accessed from http://www.crd.york.ac.uk/PROSPERO/display_record.php?

2.2. Eligibility criteria

We included studies that reported on any type of education and training intervention for recruiters to trials within any health care field, compared with no education and training, or an alternative education or training intervention. For the purpose of this review, education and training interventions are defined as structured training delivered in any format, of any duration, and using any approach, such as face to face, online, seminars, lectures, and workshops. Participants were individuals involved in recruitment to trials. This could include research nurses, general practitioners, members of the trial team, or any other individual involved in recruiting trial participants. We included only randomized trials (including cluster trials) and non-randomized (i.e., quasi-) controlled trials. We defined non-randomized controlled trials as trials where participants were allocated to the different groups using a method that was not random [8].

2.3. Outcome measures

2.3.1. Primary outcome

- Recruitment rates: proportions of eligible participants or centers recruited to the host trial. The host trial refers to any trial in which the participants of the education and training intervention were involved in recruiting individuals to.

2.3.2. Secondary outcomes

- Quality of informed consent reported by participants of either the host trial or participants of the education and training trial;
- Recruiter self-confidence;
- Host trial participants' understanding/knowledge of trial information;
- Numbers of potential trial participants approached in the host trial;
- Participants' satisfaction with training in the education/training trial;
- Retention rates to the host trial.

2.4. Search strategy and selection

We searched the following electronic bibliographic databases: EMBASE, MEDLINE, and the Cochrane Library from July 2015 (end search date of Townsend et al. [7] review) to September 2018 (date the searches were implemented). We used broad search terms such as recruitment, training, education, randomized control trials, and variations of these terms/synonyms with Boolean operators, adapted across databases (see [Appendix A](#)). No language restrictions were applied to the search strategy; however, the inclusion of studies was restricted to English language publications.

References were initially uploaded to Endnote, and duplicate citations removed. The systematic review management software, Covidence, was used for the screening process. All titles and abstracts were screened for relevance independently by at least two from a team of seven reviewers (P.C., H.D., A.H., M.H., L.M., A.P., and V.S.). Reports of studies were assessed for full-text review against the review's inclusion and exclusion criteria. Potentially relevant full-texts were uploaded to Covidence and assessed independently for inclusion by two reviewers (V.S. and H.D.). Any conflicts in decisions were discussed until agreement was achieved.

2.5. Data collection and data items

A pre-piloted data extraction form was used to extract details including study setting and aim, details of the training interventions and control conditions, numbers participating, recruitment strategies, study methodology, outcomes measured, and results. Missing data were requested from study authors as necessary. Two reviewers

(H.D. and V.S.) extracted data independently, with any discrepancies resolved through discussion and consensus.

2.6. Risk of bias

Included trials were assessed independently by a pair of reviewers (H.D. and V.S.) for methodological quality using the Cochrane "Risk of Bias" tool [9]. Studies were judged to be of low, unclear, or high risk of bias, on selection bias, performance bias, detection bias, attrition bias, reporting bias, and any other biases. Any differences between reviewers were resolved by discussion and consensus.

The unit of analysis was trial participants, for both the education/training embedded trial and the host trial. For dichotomous outcomes such as numbers recruited to the host trial, we analyzed the data based on the number of events and the number of people assessed in the intervention and comparison groups. We used these data to calculate the risk ratio (RR) and 95% confidence interval (CI). For continuous measures, we analyzed the data based on the mean, standard deviation, and number of people in the intervention and control groups to calculate mean difference and 95% CI. If more than one study measured the same outcome using different tools, we calculated the standardized mean difference and 95% CI using the inverse variance method in RevMan [10].

3. Results

3.1. Results of the search

The search yielded 14,566 records, largely because of the broad nature of the search terms. An additional five studies were sourced for inclusion from the review of Townsend et al. [7]. Of the total records, 186 were duplicates and were removed. Of the 14,385 titles and abstracts screened, 14,366 were excluded as they did not report on a trial of an education and training intervention for recruiters to trials. This resulted in 19 records assessed at full-text level. Of these 19, eight met the inclusion criteria for our review [11–18]. On further assessment, two of the eight were subsequently excluded as one was an ongoing study [18], and the second did not report on any of our pre-specified outcome measures [17]. This resulted in the inclusion of six trials in our review [11–16]. Eleven other studies assessed at full-text level were excluded [19–29] with reasons provided in [Fig. 1](#). (see [Fig. 1](#) for the search and selection flow diagram [30]).

3.2. Characteristics of included studies

[Appendix B](#) presents the summary characteristics of the included studies. Of the six included studies, two were multinational studies [14,15], two were conducted in the United States [11,13], one was conducted in the UK [16], and one in Finland [12]. Of the six included studies, four

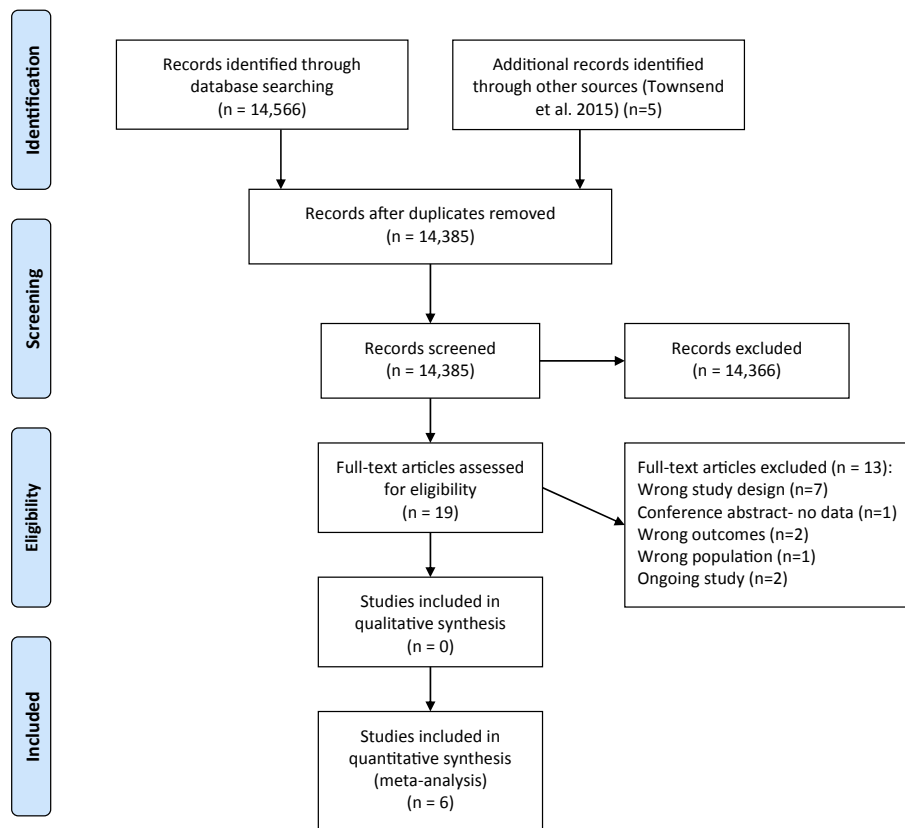


Fig. 1. Search and selection flow diagram.

[11,12,14,16] were randomized trials (of which three were cluster-randomized [11,12,16]) and two were non-randomized trials [13,15]. The education and training interventions for the included studies were implemented in the health care fields of oncology ($n = 3$), cardiovascular care ($n = 2$), and pediatrics ($n = 1$). In two of the studies, the education and training interventions were targeted at recruitment to a specific host trial, whereas three focused more generally on institutions/centers that were running trials (the remaining study did not report on the timing of the intervention). Two studies focused on the recruitment of specific groups (children and the elderly). All of the trials compared the education and training intervention with no education and training (for one study, the control group received standard information relating to recruitment, but no extra training or education [11]).

The education and training interventions in three of the studies focused mainly on communication skills [12–14]. One involved a communication skills course, lasting one evening and one morning, to improve quality of informed consent, with role play and feedback [12]. A second involved “Informed Consent Seminars” [13], and the third involved a workshop on patient information delivery and strategies to improve shared decision-making with potential trial participants [14].

Kimmick et al. [11] assessed an educational symposium along with the provision of educational materials such as

monthly mailings and lists of available protocols for use on patient charts. Kendall et al. [15] evaluated a targeted educational approach, with regular visits to the host trial sites to educate investigators and site personnel, and to help overcome any recruitment challenges. The remaining included study focused on practical ways to improve recruitment to the host trial with the provision of software for each site, and educational instructions on how to extract from data lists of patients who were potentially eligible for trial recruitment [16].

One study did not report on the number of sites included in their trial [14]. The other five trials included a total of 1,201 trial sites/institutions/centers. Three studies reported specifically on the number of individual participants enrolled to the study (both intervention and control): 132 physicians/oncologists in total (another study reported on the number of physicians attending each element of the training). Two studies reported on the number of patients involved in the host trials, which was 347 in total.

3.3. Risk of bias

Most studies were assessed overall as low or unclear risk of bias (see Figs. 2 and 3 and Appendix C). One study had low risk for allocation concealment and sequence generation, as a programmer independent of the trial created the computer-generated random allocation sequence [16].

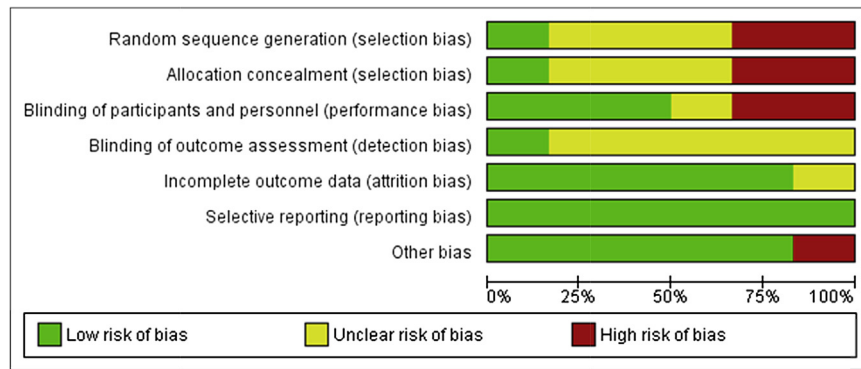


Fig. 2. Risk of bias graph.

Kendall et al. [15] and Yap et al. [13] were assessed as high risk because of non-randomization, and the remaining three studies were judged unclear because of insufficient information in the trial report to adequately assess. Owing to the nature of the interventions, it was impossible to blind participants and personnel. However, trials reporting objective measures only were assessed as low risk of bias [11,15,16], and trials reporting subjective measures only were assessed as high risk of bias [12,13].

When insufficient information was provided to assess risk of bias, we judged these to be unclear [14]. Detection bias was unclear for all studies, except that of Maxwell et al. [16], which had low risk. All studies had low attrition and reporting bias, except that of Yap et al. [13], which was

judged unclear on attrition bias. One study was judged high risk for other bias because of differences in characteristics between the control and intervention groups [12].

3.4. Effectiveness of the interventions

None of the included studies reported on the outcomes: retention rates and numbers of potential trial participants approached.

3.4.1. Recruitment rates

Three studies reported on recruitment rates. Maxwell et al. [16] reported no significant change in the cumulative randomization total over time between the training intervention and the control sites (adjusted RR 1.06, 95% CI 0.55 to 2.03, $P = 0.87$, after adjusting for site, site location, and time since the start of the intervention). Kimmick et al. [11] reported no differences in recruitment rates between training intervention and control institutions. Before the intervention, 40% of patients in intervention institutions compared with 36% in controls were recruited. During the first and second years after intervention, 36% and 31% were registered for trials in the intervention group and 32% and 31% in the control group. Kendall et al. [15] also reported on recruitment rates, and categorized sites as low- and high-recruiting sites (1–5, 6–10, 11–15, 16–20, and >20 patients) and reported that the intervention institutions had a higher proportion of high-recruiting sites; the control institutions had a higher proportion of low-recruiting sites.

3.4.2. Quality of informed consent

Quality of informed consent was reported in one study [12], using the Quality of Informed Consent Questionnaire. Significantly more patients with doctors in the training intervention group reported that “the physician offered other therapeutic options” compared with the control group (97% vs. 91%; RR 1.10, 95% CI 1.04 to 1.17) and reported a greater awareness of the study aim (89% vs. 71%; RR 1.15, 95% CI 1.03 to 1.28).

	Random sequence generation (selection bias)	Allocation concealment (selection bias)	Blinding of participants and personnel (performance bias)	Blinding of outcome assessment (detection bias)	Incomplete outcome data (attrition bias)	Selective reporting (reporting bias)	Other bias
Butow et al. 2015	?	?	?	?	+	+	+
Hietanen et al. 2007	?	?	+	?	+	+	+
Kendall et al. 2012	+	+	+	?	+	+	+
Kimmick et al. 2005	?	?	+	?	+	+	+
Maxwell et al. 2017	+	+	+	+	+	+	+
Yap et al. 2009	+	+	+	?	?	+	+

Fig. 3. Risk of bias summary.

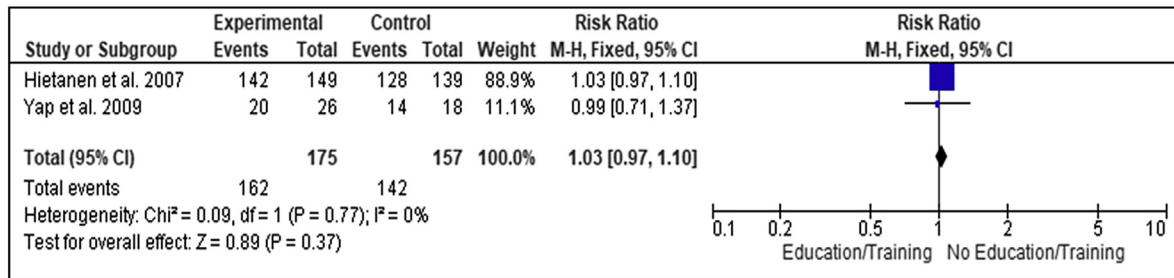


Fig. 4. Host trial participant's knowledge/understanding of trial information.

Two studies reported on the host trial participants' understanding/knowledge of trial information, including all aspects of the trial [12] or randomization only [13]. The results demonstrated no differences between the groups in understanding/knowledge of trial information (RR 1.03, 95% CI 0.97 to 1.10, 2 studies, 332 participants) (Fig. 4). Furthermore, no difference between groups was found in participants' ability to understand their voluntary choice about trial participation (RR 1.38, 95% CI 0.98 to 1.93, 1 trial, 59 participants).

3.4.3. Recruiter self-confidence

Butow et al. [14] reported narratively on recruiter self-confidence in their information provision, describing no difference from pre- to post-randomization between the training intervention and control groups.

3.4.4. Participants' satisfaction with the training intervention

Butow et al. [14] also reported on participants' satisfaction with the training intervention, describing median satisfaction scores of 57.5 (range = 41–57) and 56.0 (range = 38–73) for SGA centers (Swiss, German, Austrian) and ANZ centers (Australian/New Zealand), respectively.

4. Discussion

4.1. Summary of principal findings

This review examined the effectiveness of education and training interventions for recruiters to trials. Six trials evaluating education and training interventions were identified and included. Owing to limited evidence and differences between the included studies, in terms of outcomes and approaches, it is difficult to draw definite conclusions.

Meta-analysis was possible for one outcome, that is, host trial participant's understanding/knowledge of trial information; no differences between the groups were found. The results of the remaining reported outcomes, due to heterogeneity across the included studies, were summarized narratively. For our primary outcome of recruitment rates, reported in three of the six included studies, the evidence

of effect of education and training interventions remains conflicting. An education and training intervention, however, was found to yield greater quality in informed consent.

4.2. Strengths and limitations

This review provides contemporary, up-to-date evidence of effect underpinned by rigorous systematic review methods. Although language restrictions were not applied to the search strategy, inclusion of studies was restricted to English language publications only. This has the potential to introduce language bias and limit the scope of a review; however, none of the retrieved studies that were screened at full-text level were excluded on the basis of language, thus minimizing the potential for any language bias. We also acknowledge that identifying trials for inclusion pre-2015 was based on the Townsend et al. [7] search strategy. Although we are confident that all pre-2015 trials were captured with this search, we recognize this also as a potential limitation.

4.3. Comparison with existing literature

Similar to Townsend et al. [7], we found minimal evidence of effect of training interventions on recruitment rates to trials. Since 2015, the publication date of the Townsend review, only one further study (and one ongoing study) was identified on the effectiveness of education and training interventions. Our review thus further emphasizes Townsend's suggestion that not only are further training interventions for trial recruiters required but that these interventions should be evaluated using more robust methods to better assess impact on recruitment rates. In addition, Treweek et al. [1] highlight that further high-certainty and in-depth evidence is required around strategies to improve recruitment to trials; in particular, replication of evaluations is required to strengthen the evidence base. To add to this, we found that there has been very little evaluation, by means of controlled trials, carried out on education and training interventions for recruiters to trials specifically.

4.4. Implications for research and recruitment practice

Owing to limited evidence, it is difficult to recommend any meaningful suggestions for improving recruitment

practices. To address recruitment issues relating to trials, further work on developing a substantial evidence base around the effectiveness of education and training interventions for recruiters to trials is required. Furthermore, owing to the differences in outcomes and evaluation methods used in the included studies, it was difficult to undertake higher level syntheses such as meta-analysis on all reported outcomes, except host trial participant's knowledge/understanding of trial information. For this reason, there is a need for a standardized or core set of outcome measures for use in future trials of recruitment training interventions. The use of a core outcome set would strengthen the evidence base as it would enable enhanced comparisons between studies, and ensure that outcomes that are assessed are of relevance to key stakeholders.

5. Conclusion

There is limited evidence of effect on the impact of education and training interventions for recruiters to trials, on trial recruitment rates, so it is difficult to draw definite conclusions. To address recruitment issues relating to trials, further work on developing a substantial evidence base around the effectiveness of education and training interventions for recruiters to trials is required. In addition, there is a need for a standardized set of outcome measures, for use in future trials and systematic reviews of recruitment education and training interventions.

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Supplementary data

Supplementary data to this article can be found online at <https://doi.org/10.1016/j.jclinepi.2019.05.013>.

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