Analysis of reporting completeness in exercise cancer trials: a systematic review

CURRENT STATUS: ACCEPTED

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DOI:
10.21203/rs.2.12963/v1

SUBJECT AREAS
Health Economics & Outcomes Research

KEYWORDS
exercise, cancer, reporting, systematic review
Abstract

Background

Exercise is an effective therapeutic intervention for cancer survivors. Concerns about the completeness of reporting of exercise interventions have been raised in the literature, but without any formal analysis. This study aimed to evaluate the completeness of reporting of exercise interventions for cancer survivors in a representative sample of randomized clinical trials (RCTs).

Methods

We developed a pre-defined protocol. We searched MEDLINE, EMBASE, and CENTRAL for exercise trials in oncology between 2010 and 2017. Pairs of independent researchers screened the records, extracted study characteristics, and TIDieR criteria. The 12-item TIDieR checklist was used to generate a score of reporting completeness, based on how much of the recommended content was reported in the included trials. Because Item 5 on the checklist has two components and Item 8 has four components, the derived score was calculated as how many of the 16 components were reported, converted to a percentage.

Results

We included 131 RCTs reporting 138 interventions in the analysis. Breast cancer was the most common type of cancer (69, 50%), and aerobic exercise was the most studied exercise modality (43, 30%) followed by combined aerobic and resistance training (40, 28%). Completeness of reporting ranged from 42% to 96% among the TIDieR items; none of the items was fully reported. ‘Intervention length’ was the most reported item across interventions (133, 96%), followed by ‘rationale’ (131, 95%), whereas ‘provider’ (58, 42%) and ‘how well (planned)’ (63, 46%) were the two least reported items. Half of the TIDieR items were completely reported in 50% to 70% of the interventions, and only four items
were reported in more than 80% of the interventions (Items 2 and 8a to c). The seven items deemed to be core for replication (Items 3 to 9) exhibited a mean reporting of 71%, ranging from 42% to 96%.

Conclusion

Exercise training interventions for cancer survivors are incompletely reported across RCTs published between 2010 and 2017. The reporting of information about the provider, materials, and modifications require urgent improvements. Stronger reporting will enhance usability of trial reports by both healthcare providers and survivors, and will help to reduce research waste.

Background

Exercise is widely recognized as one of the most effective non-pharmacological interventions for improving outcomes for cancer survivors (1). A strong body of evidence suggests that cancer survivors who exercise gain benefits in quality of life (1-3), fatigue (4;5), mobility (6), depression (5), post-operative outcomes (7), and the tumor microenvironment (8;9). The first report of the American Cancer Society about exercise and cancer was published in 2003 (10); and since then, the number of randomized clinical trials (RCTs) addressing the effects of exercise in cancer survivors has grown exponentially (11).

The rapid accumulation of RCTs of exercise in cancer survivors should improve clinical outcomes, but only if the exercise interventions are reported thoroughly. Incomplete reporting of the exercise interventions impedes clinicians, researchers and patients when they attempt to interpret and implement the evidence (12-15). Incomplete reporting of interventions can also impair systematic reviews in several ways (16;17): trials may be erroneously included or excluded because of uncertainty about the intervention; and treatment differences may go unrecognized as a source of between-study variation in
effect estimates. By impairing systematic reviews, clinical decision-making is also affected (18) (19).

Complete reporting of interventions encompasses more than just naming or labelling the intervention and listing its main components; researchers must report also on other key features of the interventions, such as duration, intensity/dose, setting, mode of delivery, and monitoring (13;14;20;21). Reporting the rationale or framework that underlies the intervention can also be helpful to clinicians, who may need to adjust the intervention to suit the comorbidities or other characteristics of individual patients.

It is important to consider whether exercise reporting is complete in trials of exercise in cancer survivors because the completeness of reporting is generally lower in non-pharmacological than in pharmacological trials (13). In one review, only 39% of the non-pharmacological trials provided complete data for the intervention details (14). In a review of supervised exercise training in people with peripheral arterial disease, only around one-quarter of the trials described complete data for the mode of exercise, intensity of exercise, and tailoring/progression; and around one-tenth reported exercise intensity comprehensively (22). Similar findings were found on exercise-based cardiac rehabilitation trials (23). In reviews of trials of exercise in cancer survivors, various research groups have expressed concern about the description of the exercise protocols (1;24;25). However, no formal analysis has been published.

In order to assess the completeness of reporting, an acceptable tool is required. To encourage better reporting of trials, various checklists have been developed; for example the SPIRIT (Standard Protocol Items: Recommendations for Interventional Trials) statement for use at the protocol stage (26) and the CONSORT (Consolidated Standards of Reporting Trials) at the manuscript stage (27). An extension of the CONSORT Statement for randomized trials of non-pharmacologic treatments is also available (28). The TIDieR
(template for intervention description and replication) tool was published as an extension to these documents, to increase the detail reported about the intervention (29). The checklist contains 12 items: name, why, what, who provided, how, where, when and how much, tailoring, modifications, how well, adherence or fidelity (29). Authors of RCTs are encouraged to use the TIDieR checklist for reporting the interventions in enough detail to enable replication and facilitate the potential impact of their research on both health and society (29). The TIDieR checklist was used to rate the completeness of reporting of the intervention in the formal analyses cited above (22) (23).

The present study aimed to evaluate the completeness of reporting of exercise training interventions in RCTs that test exercise interventions in cancer survivors, using the TIDieR checklist to generate a measure of the completeness of reporting.

Methods

This study is reported according to the PRISMA statement (30) and the guidelines for reporting meta-epidemiological methodology research (31). We developed the protocol a priori and made it available via Open Science Framework (OSF) (https://osf.io/6ejh9/?view_only=4320d9fbe4134ca88422d1eaf3d5b44a; DOI 10.17605/OSF.IO/6EJH9). No major changes were introduced to the protocol throughout the conduct of this study.

Search strategy and screening

An information specialist designed (LN), tested and implemented a systematic search for RCTs published in the MEDLINE, EMBASE, and CENTRAL databases between 2010 and 2017. We selected 2010 because the CONSORT statement (27) was launched in 2010 (http://www.consort-statement.org). Additional file 1 presents the search strategies for MEDLINE, EMBASE and CENTRAL.

We used the management software Rayyan (32)) for independent screening of title and abstract. Pairs of researchers discussed disagreements, with resolution by an independent
third researcher where necessary.

**Selection criteria**

We included RCTs meeting the following criteria:

**Population**

The study population had to consist of adult (older than 18 years old) survivors of any type of cancer. A survivor was defined according to the Centers for Disease Control and Prevention (CDC), as anyone who has been diagnosed with cancer, from the time of diagnosis through the rest of life (33).

**Intervention**

We included RCTs evaluating the effects of exercise training interventions for cancer survivors. Exercise was defined as any body movement that increases energy expenditure and that is planned, structured, repetitive, and purposive in the sense that it aims to improve or maintain one or more components of physical fitness (i.e., cardiorespiratory endurance, muscular endurance, muscular strength, body composition, and flexibility) (34;35).

We accepted for inclusion any exercise training interventions involving different training modes, such as aerobic, resistance and flexibility training, as well as yoga, Qi-gong and Tai-Chi (3;11). Further, the exercise training interventions could be conducted in different settings (such as clinical or community) or mediums (such as water or land). Because of the review’s focus on the reporting of exercise training interventions, we excluded RCTs that evaluated recreational physical activity interventions rather than a structured exercise program, as well as trials reporting on manual therapy (e.g., joint mobilization techniques and therapeutic massage), cognitive-behavioral interventions, and mixed interventions that combined exercise with other therapeutic approaches, such as psychotherapy or diet or dietary advice/counseling. Finally, we excluded trials that
compared exercise training with pharmacological and surgical treatments.

Comparison

We included studies with non-exercise intervention comparisons (such as conventional care) or other exercise interventions (e.g., aerobic versus resistance training). Where trials compared two exercise interventions, both interventions were included in the analysis. For example, if there were five trials of exercise versus no intervention, and two trials of aerobic training versus resistance training, then the review would report on the completeness of reporting of nine exercise interventions.

Outcomes

We did not use the outcomes measured by the studies to determine eligibility because this study focused solely on the completeness of reporting of the intervention.

Study design and type of publication

The study design had to be a RCT. If there was any dispute about the eligibility of a trial’s design, we referred to the National Cancer Institute’s definition (36). Only full-text publications were included in the review. If the findings from a certain RCT were reported in two or more publications, pairs of reviewers evaluated all publications related to the study, retrieved the trial registry number if available, and included the primary/original publication, which was deemed to be the first publication of the study and the one with the most complete reporting of the exercise interventions.

Language

We considered for inclusion studies published in English, Spanish, Italian, Portuguese and Scandinavian languages.

Data extraction and management

Characteristics of the included studies

We extracted the following information: publication year, country, trial registry, study
name, original publication/companion, sample size (total analyzed), type of cancer, treatment stage, control group(s), exercise mode, length (weeks, with the minimum value reported in case of range), frequency (sessions/week), and setting.

**TIDieR checklist and calculation of completeness of reporting**

A pair of researchers from a pool of four worked independently to apply the TIDieR checklist to the included RCTs. As recommended by the TIDieR committee, the checklist is completed following the TIDieR guide (29), which contains an explanation and elaboration for each item. All the items were rated Yes/No. Only items that were clearly met were rated Yes; any that were partially met were rated No. We extracted data only from the full-text paper. Pairs of researchers discussed disagreements, with those outstanding resolved by an independent third researcher. Each item on the TIDieR checklist made up one component of the score of completeness of reporting, except for the multicomponent items. Item 5 contributed two components and Item 8 contributed four components. Therefore, the score was calculated as how many of 16 items were reported (as shown in Table 1) and converted to a percentage.

**Overall and subgroup analyses**

We calculated completeness of reporting in the RCTs as the number of the 16 criteria met, and presented these as percentages for each TIDieR item. We presented separate data for the subgroups of breast cancer and non-breast cancer trials.

**Table 1.** Final version of the TIDieR checklist used in this study (16 items)
1. **Brief name**
   Provide the name or a phrase that describes the intervention

2. **Why**
   Describe any rationale, theory, or goal of the elements essential to the intervention

3. **What (Materials)**
   Describe any physical or informational materials used in the intervention

4. **What (procedures)**
   Describe each of the procedures, activities, and/or processes used in the intervention

5. a. **Who provided (disciplinary background)**
   Describe the disciplinary background of the provider

   b. **Who provided (expertise, experience, or specific training)**
   Describe the expertise, experience, or specific training of the provider

6. **How**
   Describe the modes of delivery

7. **Where**
   Describe the type(s) of location(s) where the intervention occurred

   a. **When and how much (frequency)**
   Describe the number of times the intervention was delivered (e.g., number of sessions)

   b. **When and how much (length)**
   Describe the number of weeks/months the intervention lasted

   c. **When and how much (duration)**
   Describe the duration of each session (e.g., minutes/session)

   d. **When and how much (intensity)**
   Describe the intensity at which the exercise was practiced

9. **Tailoring**
   If the intervention was planned to be personalized, titrated or adapted, then describe what, why, when, and how

10. **Modifications**
    If the intervention was modified during the course of the study, describe the changes

11. **How well (planned)**
    If intervention adherence or fidelity was assessed, describe how and by whom, and if any strategies were used to maintain or improve fidelity, describe them.

12. **How well (actual)**
    If intervention adherence or fidelity was assessed, describe the extent to which the intervention was delivered as planned.

### Results

#### Results of the search

The systematic searches yielded 10702 records, and we identified 28 additional records by scrutinizing previous systematic reviews in this field. We exported 6862 records to Rayyan for screening at title and abstract, after which we read 328 records as full-text manuscripts. One reviewer (JME) retrieved all files. We included 131 RCTs in our analysis.

#### Characteristics of the included studies

The 131 RCTs contributed information about 138 interventions to the analysis. The characteristics of the individual included studies/interventions such as country, year of publication, sample size, type of cancer reported, treatment stage, and other in Additional file 2. Summary data are presented below.
Twenty-one trials (16%) provided study name or acronym, and forty-four (34%) reported their trial registry record/trial protocol, which we used as a guide to track companion studies. Walking interventions were classified as aerobic exercise. Hereafter we refer to interventions (rather than trials) as they represent our unit of analysis.

Overall, we report data from 38 countries. USA was the most common country across the analyzed interventions (38, 27%), followed by Australia (17, 12%), Canada (12, 9%), Germany (11, 8%), Korea (6, 4%), and Spain (5, 4%). Around half of the interventions were performed in groups of 10 to 50 participants (65, 47%), and one-third included 51 to 100 participants (47, 34%). Breast cancer was the most common type of cancer (69, 50%), followed by prostate cancer (20, 14%), mixed (more than one type) (14, 10%), and colorectal cancer (7, 5%). Most interventions were administered to people receiving anticancer treatment (71, 51.4%), followed by post-treatment administration (62, 45%) and pre-operative administration (6, 4.3%).

Interventions: exercise modalities

Aerobic exercise was the most studied exercise modality (43, 30.5%) followed by combined aerobic/resistance training (40, 28.4%). Resistance training alone and yoga accounted for around 13% of the interventions each. Other modalities comprised Qigong, aquatic exercise, football, high-intensity training, and Tai-Chi. One-third of the exercise interventions were implemented in clinics or hospitals. On average, exercise interventions lasted 14.3 weeks (standard deviation (SD) 12.9), and involved 2.8 (SD 2) sessions per week.

Completeness of reporting of the exercise training interventions

Total sample

Completeness of reporting ranged from 42% to 96% among the TIDieR items (see Figure 2). Intervention length was the most reported item across interventions (133, 96%),
followed by *study rationale* (131, 95%), whereas *provider* (58, 42%) and *how well (planned)* item (63, 46%) were the two least reported items. Half of the TIDieR items were completely reported in 50% to 70% of the interventions, and only four items were reported in more than 80% of the interventions (items 2 and 8a-c). In addition, the seven items (3-9) deemed to be core for replication (29) exhibited a mean reporting of 71%, ranging from 42% to 96%.

*Subgroup analysis: breast cancer*

As stated previously, this subgroup counted for half of the total sample in this study. However, completeness of reporting among the interventions for breast cancer patients exhibited similar results to the total sample. The largest difference was a 17% less complete reporting of item 7 (*where*) in the breast cancer subgroup than in the non-breast cancer subgroup. See Table 2.

Table 2. Completeness of reporting of the exercise interventions: total sample and subgroups
| TiDIEr item | Total sample n (%) | Breast cancer n (%) | Non-breast cancer n (%) |
|------------|--------------------|---------------------|------------------------|
| Item 1. Brief name | 76 (55%) | 36 (52%) | 40 |
| Item 2. Why | 131 (95%) | 66 (96%) | 65 |
| Item 3. What (Materials) | 81 (59%) | 40 (58%) | 41 |
| Item 4. What (procedures) | 108 (78%) | 53 (77%) | 55 |
| Item 5a. Who provided (disciplinary background) | 93 (67%) | 44 (64%) | 49 |
| Item 5b. Who provided (expertise, experience, or specific training) | 58 (42%) | 30 (44%) | 28 |
| Item 6. How | 104 (75%) | 54 (78%) | 50 |
| Item 7. Where | 92 (67%) | 40 (58%) | 52 |
| Item 8a. Frequency | 123 (89%) | 60 (87%) | 63 |
| Item 8b. Length | 133 (96%) | 66 (96%) | 67 |
| Item 8c. Duration | 120 (87%) | 63 (91%) | 57 |
| Item 8d. Intensity | 90 (65%) | 39 (56%) | 51 |
| Item 9. Tailoring | 82 (59%) | 37 (54%) | 45 |
| Item 10. Modifications | 70 (51%) | 32 (46%) | 38 |
| Item 11. How well (planned) | 63 (46%) | 30 (43%) | 33 |
| Item 12. How well (actual) | 70 (51%) | 35 (51%) | 35 |

**Subgroup analysis: non-breast cancer**

The other half of the interventions, those that involved non-breast cancer patients, comprised predominantly prostate cancer (20, 29%), followed by mixed type and lung cancers (each 14, 20%), and colorectal cancer (7, 10%). In general, this subgroup was reported in a more complete manner than the breast cancer subgroup and the total sample. Only a few items showed lower scores of complete reporting relative to the other
groups (Item 2, item 5b, item 6, and item 8c). See Table 2.

Discussion

Main findings
This study evaluated the completeness of reporting of exercise training interventions in a sample of RCTs in cancer survivors. Findings revealed none of the TIDieR items was fully reported across all the interventions in the RCTs. Intervention length and study rationale were the two most reported items. Conversely, relevant information for researchers, healthcare providers, and patients (such as the expertise, experience, or specific training of the provider) obtained the lowest score of reporting. We observed no major differences in the subgroups of breast cancer and non-breast cancer trials.

Comparison with previous studies
To our knowledge, this is the first study addressing the completeness of reporting in exercise trials involving cancer patients, by using the TIDieR checklist. Other studies have applied TIDieR to exercise trials in people with peripheral arterial disease (58 trials, reporting on 76 interventions) (22), trials on exercise-based cardiac rehabilitation (57 trials, reporting on 74 interventions) (23), and in trials of upper limb therapies for children with unilateral cerebral palsy (60 trials, reporting on 68 interventions) (37). In the field of exercise and cancer, Neil-Sztramko and collaborators have recently conducted relevant work about the reporting of the components and principles of resistance training prescription in breast cancer trials that measured physical fitness or body composition outcomes (38). That study found that no trials reported all components of the exercise prescription in the methods, or adherence to the prescribed intervention in the results. Similar findings were found in prostate cancer trials (39).

Strengths and weaknesses
The comprehensive systematic search run for this study as well as the independent and
duplicate conduct of the study selection and data extraction processes constitute methodological strengths. The large number of RCTs evaluated represent the largest study in using the TIDieR checklist to date. Moreover, the research team comprised a journal editor, healthcare providers, and experts in evidence synthesis in the area of exercise in cancer and other chronic conditions. We believe the decision of splitting the number of TIDieR items in our analysis provides a more specific insight to readers. This approach was supported by the developers of the TIDieR tool.

Despite these methodological strengths, we cannot state with certainty that our analysis covered all available RCTs in exercise and cancer; it is likely some references were not identified. Also, the inconsistencies in the trials registry records made it difficult to track companion studies. However, because we analysed a large sample consisting of all eligible trials consecutively identified within a database with extremely high coverage of the relevant trials (40;41), and two other broad-ranging databases, it is likely that the sample is highly representative of all trials of exercise in cancer.

**Implications for practice and further research**

Findings from this study encourage researchers to adhere to international reporting guidance when formulating and publishing their research in order to facilitate translations of their findings into practice. Our results indicate there is still work to be done in this regard.

Future research could examine the reporting of some additional items, some of which have more recently been listed in the Consensus on Exercise Reporting Template (CERT) tool (42). These factors include whether the exercise is supervised and whether motivational strategies are used. We did not know about the CERT tool at the time our study protocol was formulated, and decided that it was not worth changing the study protocol to incorporate the CERT a posteriori, because of the substantial overlap between it and
TIDieR.

Journals should encourage trial authors to adhere to reporting guidance when processing submissions. Thus, journals should endorse checklists for reporting interventions as they do for CONSORT or any other related statements (29). Hopewell et al. (43) found in a time series design that an active implementation of the CONSORT for abstracts guidelines by journals improved the number of checklist items reported in abstracts of randomized trials. Journals might ask researchers to use TIDieR and perhaps CERT in conjunction when completing item 5 of the CONSORT checklist, and there refer the reader to a detailed assessment of the intervention-reporting checklist.

Conclusion

Exercise training interventions for cancer survivors are reported moderately well among RCTs published between 2010 and 2017. The reporting of information about the provider, materials, and modifications requires urgent improvement. More complete reporting of exercise training interventions for cancer patients will enhance trial usability for both healthcare providers and patients, and will contribute to a large extent in the battle to reduce research waste (44;45). Researchers might use the TIDieR checklist when reporting their exercise interventions in further trials.

List Of Abbreviations

Consensus on Exercise Reporting Template (CERT); Consolidated Standards of Reporting Trials (CONSORT); Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA); Randomized Clinical Trial (RCT); Standard Protocol Items: Recommendations for Interventional Trials (SPIRIT); Template for Intervention Description and Replication (TIDieR)

Declarations
**Ethics approval and consent to participate**

Not applicable

**Consent for publication**

Not applicable

**Availability of data and material**

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

**Competing interests**

The authors declare that they have no competing interests

**Funding**

No funding was received for this study

**Authors' contributions**

JME and JB conceived the initial idea of this study. All authors help develop the study protocol. JM and ME contacted developers of the TIDieR checklist and tailored the final version of the checklist. LN developed the search strategy and search the databases. JME, JB, IR and JMT extracted data. JM analyzed the data. JME lead the writing of the study with feedback from all authors. All authors read and approved the final version of the manuscript.

**Acknowledgements**

We thank Dr. Tammy C Hoffmann for her insight and suggestions regarding the generation of a score of reporting completeness based on the TIDieR checklist.

**Authors' information (optional)**

Jose Meneses-Echavez and Julia Bidonde have large experience in evidence syntheses for exercise and chronic conditions. Mark Elkins currently serves as editor of the Journal of Physiotherapy. Indira Rodriguez and Javier Martinez have clinical experience in exercise
prescription for the management of chronic patients and analysis of big datasets.

A preliminary analysis was presented as a poster at the 25th Cochrane Colloquium 2018 in Edinburgh.

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Additional Files

Additional file 1. Search strategy

Additional file 2. Characteristics of the included studies (n = 131 RCTs; 138 exercise interventions)

Figures

![Flow diagram for the selection of the studies.](image)

Figure 1

Flow diagram for the selection of the studies.
Completeness of reporting of the exercise interventions: total sample (n=131 RCTs; 138 exercise interventions)

Supplementary Files

This is a list of supplementary files associated with the primary manuscript. Click to download.

Additional File 1.pdf
Additional File 2.pdf