Benchmarking Controlled Trial—a novel concept covering all observational effectiveness studies

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The Benchmarking Controlled Trial (BCT) is a novel concept which covers all observational studies aiming to assess effectiveness. BCTs provide evidence of the comparative effectiveness between health service providers, and of effectiveness due to particular features of the health and social care systems. BCTs complement randomized controlled trials (RCTs) as the sources of evidence on effectiveness. This paper presents a definition of the BCT; compares the position of BCTs in assessing effectiveness with that of RCTs; presents a checklist for assessing methodological validity of a BCT; and pilot-tests the checklist with BCTs published recently in the leading medical journals.

Key messages

- The Benchmarking Controlled Trial (BCT) is a novel concept which covers all observational studies aiming to assess effectiveness.
- BCTs assess difference in effectiveness between single or a set of intervention(s), between clinical pathways, or between interventions targeting health care system factors with an aim to increase effectiveness.
- Published BCTs have currently several methodological limitations, some of which could be avoided, and others should be acknowledged.
- BCTs support both clinical and policy decisions, and should be given a high priority in research and in improvement activities.

Introduction

The experimental studies, randomized controlled trials (RCTs), provide the least biased information of the efficacy of medical interventions and create the basis for systematic reviews on effectiveness of interventions (1). However, RCTs mostly assess effectiveness of interventions in ideal settings, and they focus on specific interventions rather than considering how effective is the whole clinical pathway (from the first treatment through all interventions during e.g. a 1-year follow-up time)—the latter is crucial for overall effectiveness. Thus there is a need for valid observational data on actual performance in routine settings, particularly as all educational, research, and leadership activities in medicine are intended to advance the health of the general population and care of ordinary patients (2,3).

The first aim of this paper is to assess the need for the new concept of Benchmarking Controlled Trials (BCTs), provide a definition of the BCT, and to present the two main categories (clinical, and health and social care system-related), and the respective subcategories of BCTs. The second aim is to present a checklist for assessing the methodological validity of a BCT and to point out methodological differences between RCTs and BCTs. The third aim is to pilot-test the checklist with BCTs published recently in the leading medical journals.

Methods

The previous international recommendations on how to report observational studies and systematic reviews of them (4,5) provide guidance on studies that investigate associations between exposures and health outcomes and address three types of observational studies: cohort, case-control, and cross-sectional studies. The author’s idea was that there is a need for a framework which starts from the study question of effectiveness in observational settings. When the aim is to assess effectiveness of interventions, there are two options: experimental design (randomized controlled trials) or observational design. This paper concentrates on observational designs, and presents a comprehensive framework for them within the novel concept of Benchmarking Controlled Trials (BCTs).

When assessing effectiveness in an observational (real-world) setting, the index and comparator groups must have a priori as similar groups of patients as possible in order to allow adjusting for the potential baseline incomparability. Therefore, the comparisons have to be made between peers treating similar patients and thus there is always an element of benchmarking involved. This is the reason for the concept Benchmarking Controlled Trial. In addition, using e.g. a term such as observational controlled trial
would probably have connotations that do not coincide with the present new idea (6).

Differentiating the two main BCT categories—clinical and health care system determinants for effectiveness—was based on the author's idea that the requirement for baseline comparability for the clinical comparisons is, indeed, equally much needed when studying interventions aimed to make changes in the health care system (and through these changes increase effectiveness of interventions).

The pertinent clinical subcategories were consequently: 1) effectiveness of a particular single or set of interventions during a limited time frame (like surgery, or 3 months' rehabilitation period) and 2) effectiveness of the whole clinical pathway from start (e.g. acute myocardial infarction) through all various health (and social) care interventions (diagnostic, treatment, rehabilitation; primary, secondary, tertiary care) which happen during e.g. a 1-year follow-up time. The health care system intervention subcategories were defined further according to recent literature (Figure 1) (7). For health care system interventions no universally established categories exist, but, regardless of what they are, any change in the health care system aiming to increase effectiveness falls into the category of a BCT.

The checklist for methodological validity issues of BCTs, as well as the appraisal of methodological issues inherent to BCTs, was based on the author's previous work with randomized controlled trials and observational studies (8–13), and with methodological issues in RCTs and observational effectiveness studies, including work within the Cochrane Collaboration Back Review Group (1,9,12,14–16). Previous checklists for observational studies and systematic reviews of them were also utilized (STROBE (4), MOOSE (5)), as well as scientific literature on particular characteristics of observational studies relevant in the assessment of effectiveness of interventions (17).

For piloting the checklist, the 10 most recent BCTs published in the leading medical journals (New England Journal of Medicine, Lancet, Journal of American Medical Association, British Medical Journal, and Annals of Internal Medicine) were identified through a PubMed search and by the author searching the articles directly from the journals. The search terms were: benchmarking, registries, effectiveness, and name of the journal. All the included articles had to have an observational design, and aim to assess effectiveness of an intervention directed to patients or directed to the health care system. Five articles assessing clinical features and five assessing health care system-related features as determinants of effectiveness from January 2010 to October 2014 were included. Data extraction was rechecked, and errors were corrected by the author to reach the final appraisal.

Results

Definition and categories of the Benchmarking Controlled Trial

There is a clear need for the new concept Benchmarking Controlled Trial (BCT) as there is no previous systematic guidance on methodological issues in planning and reporting an observational effectiveness study (4,5). Furthermore, the idea of the author that, in addition to clinical interventions, any intervention directed to the health care system must be studied in a BCT is a new one. The term benchmarking is accurate because all comparisons have to be between peers and thus include an element of benchmarking. Furthermore, the results of BCTs should be exploited in the effort to increase effectiveness using the comparative data between peers—which is benchmarking (2).

A BCT is defined as an observational study aiming to provide non-biased estimates of comparative differences in outcomes and costs in real-world circumstances due to a single or a set of intervention(s) or throughout the clinical pathway between two or more health service providers for a well-defined group of patients; or an observational study aiming to provide evidence of the comparative effectiveness of the health care system or parts of it among a well-defined group of patients. Data on disadvantaged
patient groups should be included always when feasible, because their prognosis often differs from that of non-disadvantaged groups. Therefore, inability to control for the differences between disadvantaged and non-disadvantaged populations may lead to biased estimates. Furthermore, data on prevailing inequality will be go unnoticed.

The study question in BCTs should ideally be defined according to the PICO principle (patient, intervention, comparison, intervention, and outcome) taking into consideration interventions during the whole clinical pathway. The health care service providers can be individuals, health care units, hospitals, health care districts, or countries.

Features of BCTs in the two main categories (clinical effectiveness and factors related to the health care system) and in their subcategories are presented in Table I. Figure 1 illustrates the categories and subcategories of BCTs covering all observational study designs on effectiveness. In order to illustrate the entity of effectiveness studies, also RCTs are shown in the picture, as well as their subcategories explanatory (ideal circumstances) and pragmatic (ordinary health care circumstances). It must be emphasized that although pragmatic RCTs provide evidence on effectiveness in routine settings, they seldom cover the whole clinical pathway, and generalizability to other settings is limited.

**Characteristics of the checklist and methodological issues in BCTs**

The main categories and their subcategories of methodological issues in BCTs are presented in Table II. The pilot-testing of the checklist shows also main contents of the 10 studies.

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**Table I. Categories, subcategories, and characteristics of Benchmarking Controlled Trials (BCTs).**

| BCT categories and subcategories | Study objective | Design issues | Causal and effect factors | Implications for |
|----------------------------------|-----------------|---------------|---------------------------|-----------------|
| 1. Clinical comparison           | To assess differences in outcome between health care providers (individual, hospital, district, country) who treat similar patients but their way of treating patients (from single intervention to clinical pathway) potentially differs | 1. Between-group differences at baseline must be adjusted for | Causal factor: differences in single or sets of interventions or in clinical pathways between the comparator arms | Clinicians, policy-makers |
| (as determinants for effectiveness and efficiency) | 2. Diagnostics and treatment procedures during the clinical pathway should be properly documented (i) to appraise how plausible the differences in outcome are, and (ii) to make decisions on how to improve treatment of patients | Effect factor: differences in all relevant outcomes between the comparator arms | Policy-makers |
| Subtypes:                        |                 |               |                           |                 |
| 1.1. single or set of intervention(s) |                 |               |                           |                 |
| 1.2. whole clinical pathway      |                 |               |                           |                 |
| 2. System comparisons of the health and social care system (as determinants for effectiveness and efficiency) | To assess differences in outcome between health care providers due to reasons related to the health and social care system | 1. Between-group differences at baseline must be adjusted for | Causal factor: differences in features related to the health care system or part of it |                 |
| Subtypes:                        |                 |               |                           |                 |
| 2.1. related to the financing of the care system (e.g. tax-based or insurance-based system) |                 |               |                           |                 |
| 2.2. related to the reimbursement and incentives (e.g. fee for service, bonus for quality) |                 |               |                           |                 |
| 2.3. related to how and by whom the services are organized/provided (e.g. centralized versus decentralized) |                 |               |                           |                 |
| 2.4. related to the regulations (e.g. on uptake of new technology) |                 |               |                           |                 |
| 2.5. related to the available resources for health care (e.g. amount of personnel, GDPs of the countries) |                 |               |                           |                 |
| 2.6. related to other system or structure-related issues (e.g. freedom of choice) |                 |               |                           |                 |

It is noteworthy that there is an overall methodological difference between experimental trials and benchmarking trials. In experimental trials (RCTs) the data collection in each treatment arm is determined in a uniform way, and researchers’ obligation is that the conduct of an RCT adheres to the protocol. In observational settings—comparing different service providers—the accrual of the data may not be determined beforehand as strictly as in an RCT, or the quality assurance during data gathering may not be as rigorous. Therefore validity assessment in BCTs must usually be undertaken separately for all the health care service providers. Even if there has been uniform instructions on how to collect the data, the success of doing so may differ between the providers.

Another notable methodological issue is that when assessing the comparative effectiveness of a particular intervention or the whole clinical pathway in BCTs, appropriate baseline adjustment is a major challenge. Obtaining proper information of the interventions during the clinical pathway is also most important for two reasons: Firstly, to get further evidence supporting the plausibility of differences in effectiveness estimates, and secondly to have information to be used for improving the treatment processes.

When assessing the effectiveness of interventions targeting the health care system there are four major challenges. Firstly, sufficient data are needed to obtain information indicating whether the health care system factors (e.g. related to an economic incentive) may have led to selection of patients and thus to differences in baseline characteristics. The second challenge is to obtain data of the patients’ clinical pathways to know in what degree the intervention targeting the system may have changed.
the way patients are treated. The third challenge is to adjust for differences in baseline characteristics between the comparators, and analyze differences in treatment processes as mediators of the effects posed by the health care system factors. The fourth challenge is to try to document all the effects the intervention causes to the health care system including unintended unfavorable effects. However, this major challenge of observing a complex system goes beyond the present treatise.

A big difference between benchmarking controlled trials (BCTs) and randomized controlled trials (RCTs) is selection of patients. In the former, patients entering the study in each treatment arm may differ due to selection, while in the latter random allocation to treatment arms (regardless of selection) leads often to comparable treatment groups. To decrease potential for selection bias in BCTs, a two-step procedure is suggested: 1) eligibility criteria should be chosen so that they lead to a homogeneous patient population (e.g. only patients having their first-time acute ischemic stroke will be included) (13,18), and 2) the residual baseline differences have to be statistically adjusted. Instrumental variables may be feasible in some cases to compensate partially for the lack of randomization (19), and the propensity score method may enhance baseline comparability in BCTs (8). Exploitation of a natural experiment may provide an excellent opportunity to increase baseline comparability in BCTs; e.g. in a previous study the health effects of becoming unemployed were studied in a situation when due to nationwide recession suddenly half of construction workers become unemployed, and the allocation to unemployment occurred mainly by chance (20).

Concerns of sufficient clinical information and validity of the data are usually greater in BCTs than in RCTs—particularly if the data for a BCT have been gathered retrospectively, and thus no a-priori protocol has been used. A high number of dropouts is a validity concern for both RCTs and BCTs, as well as the importance of using valid outcome measures. Selective outcome reporting by researchers within a RCT may lead to biased conclusions, but in BCTs selective reporting may occur also during the data collection—often undertaken by the health care providers themselves. There are a number of statistical analysis issues that are characteristic to BCTs (Table II).

**Pilot-testing of the checklist**

All the 10 articles were from the *New England Journal of Medicine* and *Lancet*, as eligible studies were not found from the other journals (Table II) (21–30).

In the five studies assessing clinical effectiveness, the diagnoses included treatments for selected cancers, non-cardiac surgery, bariatric surgery, rupture of an aortic aneurysm, and acute myocardial infarction. The main outcomes were mortality in four studies, and complication rates in one study. In the five studies assessing effectiveness in relation to health care system-related factors, the indications were more varied than in the clinical effectiveness studies and included a set of surgical indications (two BCTs), a set of indications treated conservatively, intensive care patients, and ambulatory care patients. The determinants for the outcome were the size of the centers providing the service, quality improvement program, presence of a night-time intensivist in the hospital, pay-for-performance, and workload and qualifications of nurses. The main outcomes were mortality in four studies, and health care spending and quality of care in one study.

Concerning methodological issues in the 10 studies several limitations were observed. No study provided a description of patients’ clinical path prior to eligibility for the study. No study exploited an opportunity provided by a natural experiment. Valid diagnostic information at baseline was presented by four studies with a clinical research question, and in two of the studies with a health care system-related objective. There were deficiencies in other clinical baseline factors; and factors indicating lifestyle or environment were lacking in all the studies. Information of diagnostics and treatment procedures was lacking altogether in one clinical study and in three studies with focus on the health care system. No study assessed outcomes among disadvantaged patient groups. No study utilized instrumental variables, and only two studies provided power calculations for determining size of the study sample.

**Discussion**

This paper presents a novel concept, the Benchmarking Controlled Trial (BCT). There are several new ideas involved, particularly 1) that an element of benchmarking is always involved when making observational comparisons in real-world circumstances, and 2) that assessment of effectiveness due to any health care system intervention faces the same methodological challenges as clinical comparisons. Because of the risk for more than one connotation for one concept, a new term of e.g. observational controlled trial did not seem to be appropriate (6).

In those BCTs which pursue evidence on clinical effectiveness, information of baseline patient characteristics, of diagnostic procedures and treatments, and of the outcomes is needed for the comparisons between providers. If baseline imbalances between patients treated by different providers can be satisfactorily adjusted for, also comparisons based on treatment outcomes may be justified (31). If feasible, all clinically important patient-relevant outcomes should be documented. However, it is most important to obtain data also of the treatment processes—how well these concord with current scientific evidence (32). Benchmarking controlled trials should aim to assess quality (appropriate interventions), effectiveness and costs of services, as well as issues related to potential inequality in obtaining services shown effective (3).

In BCTs which pursue evidence on effectiveness due to health care system-related factors, there must be a homogeneous target population, and if there are several diagnoses, they should preferentially be differentiated and evidence presented separately for each diagnosis. If there is insufficient data of the diagnoses and related baseline characteristics, the evidence on effectiveness may remain very uncertain.

Previous checklists for advancement of better reporting of observational studies give guidance for studies aiming to assess causal relationship between exposure and outcome. The checklist developed for and described in this paper is intended for supporting planning, conducting, reporting, and peer reviewing manuscripts of observational studies assessing effectiveness of interventions, the BCTs.

The pilot-testing of the checklist using recent articles published in leading medical journals showed a wide variety of methodological strengths and limitations in the original studies. No study provided a description of patients’ clinical path before entering the study. Description of baseline characteristics was deficient or even lacking, causing uncertainty in between-group comparability. Information of diagnostics and treatment procedures was scarce. Instrumental variables were not utilized, and power calculations were rare.

**Conclusions**

The new concept of the BCT provides guidance for studies assessing comparative effectiveness between single or sets
### Table II. Methodological characteristics of Benchmarking Controlled Trials (BCTs) in 10 studies published between January 2010 and October 2014 in leading medical journals. Assessment is based solely on each particular paper; if information is not reported, the issue is assessed as unclear. Each characteristic is recorded as yes, partial, unclear, or no; yes indicates that the criterion has been met.

| Study characteristics | Coleman et al., Lancet, 8 Jan 2011* | Pearse et al., Lancet, 22 Sep 2012 | Birkmeyer et al., NEJM, 10 Oct 2013 | Karthikesaliam et al., Lancet, 15 Mar 2014 |
|-----------------------|--------------------------------------|---------------------------------|-----------------------------------|------------------------------------------|
| 1. Research question and study design | To produce up-to-date survival estimates for selected cancers, to establish whether international differences (Australia, Canada, Denmark, Norway, Sweden, UK) in survival have changed, and to investigate the causes of survival deficits | Describe mortality rates and patterns of critical care resource use for patients undergoing non-cardiac surgery across several European nations | To assess the effect of surgical skill as a determinant for complication rates after bariatric surgery | To compare the in-hospital mortality of patients with rupture of an abdominal aortic aneurysm in England and USA |
| 1.1. clinical or system comparison | Clinical | Clinical | Clinical | Clinical |
| 1.2. subcategory of comparison | Whole clinical pathway | Whole clinical pathway | Single intervention | Whole clinical pathway |
| 1.3. conceptually pertinent and clear | Yes | Yes | Yes | Yes |
| 1.4. natural experiment (allocation to study groups apparently by chance) | No | No | No | No |
| 1.5. operationalized according to the PICO principle (patient; treatment; comparison treatment; outcomes) | No | No | Yes | Yes |
| 2. Selection of patients/population to the study and measures to increase comparability (all studies have individual patient data) | Population-based register | Clinical sample | Clinical register | Administrative databases |
| 2.1. population-based cohort, administrative database, or clinical register | Retrospective Country level | Prospective Country level | Prospective Individual provider | Unclear Country level |
| 2.2. prospective or retrospective design | No | No | NA | No |
| 2.3. level of health care provider (e.g. individual, health care center, hospital, district, country) | Yes | No | Yes | Yes |
| 2.4. description of patients' clinical path before eligible for the study | Yes | No | No | No |
| 2.5. description of patients' clinical eligibility criteria | Yes | Yes | No | Yes |
| 2.6. comprehensive patient population of the catchment area | No | No | No | No |
| 2.7. restriction of patients to a particular group in order to increase homogeneity (e.g. first episode ever of ischemic stroke) | No | No | No | No |
| 2.8. use of instrumental variables to compensate for lack of randomization | No | No | No | No |
| 3. Validity and completeness of baseline data + Comparability ensured between groups at baseline (e.g. Validity: Yes; Comparability: No → Yes/No) | Yes/Yes | Yes/Yes | Yes/Yes | Yes/Yes |
| 3.1. diagnostics | Yes/Unclear | No/Unclear | Yes/Unclear | Yes/Unclear |
| 3.2. other clinically important data relevant to the particular disorder/disease (e.g. severity) | No/Unclear | No/Unclear | No/Unclear | No/Unclear |
| 3.3. general health/risk status | No/Unclear | No/Unclear | No/Unclear | No/Unclear |
| 3.4. co-morbid conditions | No/Unclear | No/Unclear | No/Unclear | No/Unclear |
| 3.5. behavioral factors (e.g. on health-related lifestyle) | No/Unclear | No/Unclear | No/Unclear | No/Unclear |
| 3.6. environmental factors (e.g. work conditions) | No/Unclear | No/Unclear | No/Unclear | No/Unclear |
| 3.7. potential inequality (e.g. socioeconomic status) | No/Unclear | No/Unclear | No/Unclear | No/Unclear |
| 3.8. other potential predictors (e.g. genetic factors), confounders, and effect modifiers | No/Unclear | No/Unclear | No/Unclear | Unclear/Unclear |
| 4. Validity and completeness of process data (also unrelated to the disorder in question) throughout the clinical pathway | Yes | Yes | Yes | Yes |
| 4.1. diagnostics | | | | |
| 4.2. treatment procedures | No | No | Yes | Yes |
To assess whether differences in patient-to-nurse workloads and nurses’ educational qualifications in nine countries with similar patient discharge data are associated with variation in hospital mortality after common surgical procedures.

### Table II. Methodological characteristics of Benchmarking Controlled Trials (BCTs) in 10 studies published between January 2010 and October 2014 in leading medical journals.

| Study                                      | Clinical register | Administrative databases | Clinical register | Clinical register | Clinical register | Administrative databases |
|--------------------------------------------|-------------------|--------------------------|-------------------|-------------------|-------------------|--------------------------|
| Chung et al., NEJM, 2 June 2011            | Yes               | No                       | No                | No                | No                | No                       |
| Finks et al., NEJM, 8 Nov 2012a           | No                | No                       | No                | No                | No                | No                       |
| Song et al., NEJM, 31 May 2012b           | Partial           | Unclear                  | No                | No                | No                | No                       |
| Wallace et al., NEJM, 9 Aug 2011           | Unclear           | Yes                      | No                | No                | Yes               | No                       |
| Sutton et al., NEJM, 15 Mar 2014           | Yes               | No                       | No                | No                | Yes               | No                       |
| Aiken et al., Lancet, 24 May 2014          | Yes               | Yes                      | No                | No                | No                | No                       |

(Continued)
### Table II. (Continued)

| Study characteristics                                                                 | Coleman et al., Lancet, 8 Jan 2011<sup>a</sup> | Pearse et al., Lancet, 22 Sep 2012 | Birkmeyer et al., NEJM, 10 Oct 2013 | Karthikesalinam et al., Lancet, 15 Mar 2014 |
|---------------------------------------------------------------------------------------|-----------------------------------------------|-----------------------------------|-------------------------------------|---------------------------------------------|
| 4.3. rehabilitation                                                                    | No                                            | No                                | NA                                  | NA                                          |
| 4.4. hospitalizations and health care visits                                          | No                                            | No                                | Yes                                 | Yes                                         |
| 4.5. individual behavior (e.g. lifestyle-related to health)                           | No                                            | No                                | No                                  | NA                                          |
| 4.6. adherence to treatments                                                           | Yes                                           | No                                | Yes                                 | NA                                          |
| 4.7. characteristics of the clinical pathway                                          | No                                            | No                                | NA                                  | NA                                          |

5. Validity and completeness of outcome data (related to the disorder in question)

| 5.1. validity of the outcomes                                                          | Yes                                           | Yes                                | Yes                                 | Yes                                         |
| 5.2. outcomes assessed also among disadvantaged patients                               | No                                            | No                                | No                                  | No                                          |
| 5.3. comparability (similarity) of follow-up time points                               | Yes                                           | Yes                                | Yes                                 | Yes                                         |
| 5.4. percentage of dropouts during follow-up documented and acceptable (< e.g. 10%)   | Yes                                           | Yes                                | Yes                                 | Yes                                         |
| 5.5. data at each comparator arm free of suggestion of selective outcome reporting     | Yes                                           | Yes                                | Yes                                 | Yes                                         |

6. Statistical and data issues

| 6.1. description of power calculations and rationale on how the study size was arrived at or post-analysis power calculation | No                                            | Yes                                | No                                  | No                                          |
| 6.2. documentation of how data were classified and coded (e.g. blinding, multiple raters, inter-rater reliability)         | Unclear                                       | Unclear                           | Yes                                 | Unclear                                     |
| 6.3. measures to increase reliability of data classification and coding (e.g. blinding, multiple raters, inter-rater reliability) | Yes                                           | Unclear                           | Yes                                 | Unclear                                     |
| 6.4. description of all primary statistical methods, including those used to control for confounding | Yes                                           | Yes                                | Yes                                 | Yes                                         |
| 6.5. description and use of methods to examine subgroups and interactions              | No                                            | Yes                                | Yes                                 | Yes                                         |
| 6.6. description and use of propensity score or other methods to improve comparability at baseline | No                                            | No                                | No                                  | No                                          |
| 6.7. adjustment for the characteristic outcomes of each health care provider (e.g. differences in general life expectancy in each country) | Yes                                           | No                                | NA                                  | No                                          |
| 6.8. incomplete outcome data adequately addressed. If no missing data or appropriate imputation: Yes                         | Yes                                           | Yes                                | Yes                                 | Yes                                         |
| 6.9. use of multilevel modeling or survival modeling                                  | Yes                                           | Yes                                | Yes                                 | Yes                                         |

In studies on health care system as determinant of outcome:

| 6.10. diagnostic and treatment processes analyzed as the mediators of the effects     | No                                            | No                                | No                                  | No                                          |

Study having comparisons also with cohorts in time, i.e. changes in outcomes between follow-up years (each have patients of their own).

For this design there are three additional methodological issues:

| 7.1. Documentation of changes in patient characteristics over time                   | No                                            |                                    |                                      |                                             |
| 7.2. Documentation of changes in treatment practices over time                       | No                                            |                                    |                                      |                                             |
| 7.3. Documentation of changes in patient outcomes over time                          | Yes                                           |                                    |                                      |                                             |

NA = not applicable.

<sup>a</sup>Study having comparisons also with cohorts in time: study characteristics 7.1–7.3.

<sup>b</sup>Study assessing the effect of one factor related to the organization of the system (e.g. presence of night-time intensivist).
| Study                          | No | Yes | NA | No | No | No | No | No | No | No | No | No |
|-------------------------------|----|-----|----|----|----|----|----|----|----|----|----|----|
| Chung et al., Lancet, 12 April 2014 | No | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes |
| Finks et al., NEJM, 2 June 2011 | No | No | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| Song et al., NEJM, 9 Aug 2011  | No | No | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| Wallace et al., NEJM, 31 May 2012 | No | No | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| Sutton et al., NEJM, 8 Nov 2012 | No | No | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |
| Aiken et al., Lancet, 24 May 2014 | No | No | NA | NA | NA | NA | NA | NA | NA | NA | NA | NA |

- **4.3. rehabilitation**: No No NA NA Y es No NA NA No No
- **4.4. hospitalizations and health care visits**: No No Y es Y es NA No NA NA No No
- **4.5. individual behavior (e.g. lifestyle-related to health)**: No No No NA No No NA NA No No
- **4.6. adherence to treatments**: Y es No Y es NA No No NA NA No No
- **4.7. characteristics of the clinical pathway**: No No NA NA No No NA NA No No
- **5. Validity and completeness of outcome data (related to the disorder in question)**
  - **5.1. validity of the outcomes**: Y es Y es Y es Y es Y es Y es Y es Y es Y es Y es
  - **5.2. outcomes assessed also among disadvantaged patients**: No No No No No No No No No No
  - **5.3. comparability (similarity) of follow-up time points**: Y es Y es Y es Y es Y es Y es Y es No Y es Y es
  - **5.4. percentage of dropouts during follow-up documented and acceptable**: Y es Y es Y es Y es Y es Y es Y es Y es Y es Y es
  - **5.5. data at each comparator arm free of suggestion of selective outcome reporting**: Y es Y es Y es Y es Y es Y es Y es Y es Y es Y es
- **6. Statistical and data issues**
  - **6.1. description of power calculations and rationale on how the study size was arrived at or post-analysis power calculation**: No Y es No No No No Y es No No No
  - **6.2. documentation of how data were classified and coded (e.g. blinding, multiple raters, inter-rater reliability)**: Unclear Unclear Y es Unclear Y es Unclear Unclear Y es Unclear Unclear
  - **6.3. measures to increase reliability of data classification and coding (e.g. blinding, multiple raters, inter-rater reliability)**: Y es Unclear Y es Unclear Partial Unclear Unclear Y es Unclear Unclear
  - **6.4. description of all primary statistical methods, including those used to control for confounding**: Y es Y es Y es Y es Y es Y es Y es Y es Y es Y es
  - **6.5. description and use of methods to examine subgroups and interactions**: No Y es Y es Y es Y es Y es Y es Y es Y es Y es
  - **6.6. description and use of propensity score or other methods to improve comparability at baseline**: No No No No Y es No Y es No No No
  - **6.7. adjustment for the characteristic outcomes of each health care provider (e.g. differences in general life expectancy in each country)**: Y es No NA No No No No No No No
  - **6.8. incomplete outcome data adequately addressed. If no missing data or appropriate imputation**: Y es Y es Y es Y es Y es Y es Y es Y es Y es Y es
  - **6.9. use of multilevel modeling or survival modeling**: Y es Y es Y es Y es Unclear Y es Y es No No Y es

In studies on health care system as determinant of outcome:

- **6.10. diagnostic and treatment processes analyzed as the mediators of the effects**: No No No No No Y es No No No No

For this design there are three additional methodological issues:

- **7.1. Documentation of changes in patient characteristics over time**: No No
- **7.2. Documentation of changes in treatment practices over time**: No No
- **7.3. Documentation of changes in patient outcomes over time**: Y es Y es

NA = not applicable.

a Study having comparisons also with cohorts in time: study characteristics 7.1.–7.3.

b Study assessing the effect of one factor related to the organization of the system (e.g. presence of night-time intensivist).
of interventions, between clinical pathways, or between health care systems or factors related to the system. Benchmarking controlled trials cover the whole area of observational effectiveness research.

A checklist for assessing the methodological validity of BCTs has here been subjected to preliminary pilot-testing, but should be properly validated. However, the checklist can readily be used in planning, conducting, reporting, and appraising BCTs.

Current BCTs seem to have several methodological limitations, some of which could be avoided in planning and conducting phases of the studies, and others should be acknowledged in discussion.

Benchmarking controlled trials—supporting both clinical and policy decisions—should be given a high priority in research, and their results should be used in improvement activities provided they have sufficient methodological rigor and generalizability. The proposed methodology is suggested also for non-scientific quality improvement and benchmarking undertakings.

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