How to increase the value of randomised trials in COPD research

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ABSTRACT: Methodological criteria that increase the validity of randomised trials are often not considered in respiratory research, even in large chronic obstructive pulmonary disease (COPD) trials. We describe four important aspects in the design, analysis and reporting of randomised trials, selected based on their relevance to current COPD research and based on our judgments of importance for researchers and users of the literature.

First, to optimally control for confounding, where confounding refers to a factor that is associated with an exposure or intervention and influences the outcome, a clear definition of the main relationship between treatment and the primary outcome as well as identification of measurable confounders is required. In addition to randomisation per se as the key method to protect against confounding, restriction (excluding patients with specific characteristics that may introduce confounding), stratification (separate randomisation of patients with specific characteristics) and statistical adjustment are means to be considered to optimally control for confounding that simple randomisation may not achieve.

Secondly, the selection of the primary outcome should be guided by the importance to patients. Secondary outcomes provide hypotheses about the effects observed for the primary outcome and can provide important data for systematic reviews and meta-analyses, but should be interpreted with caution in single trials.

Thirdly, in study power calculations, not only the actual sample size, but the number of events, has a large influence on the power of the study and, often, unrealistic assumptions about event rates are made to increase the feasibility of trials.

Finally, essential steps to transfer results from research to practice include complete reporting of trials and developing tools, such as decision aids, to support patients and physicians in their shared decision making.

KEYWORDS: Chronic obstructive pulmonary disease, evidence-based medicine, study design
application of trials [4]. The aim of this article is, therefore, to
discuss selected methodological issues in the design, analysis
and interpretation of randomised trials that inform readers and
inform investigators of future studies. We selected the
following four issues based on the limitations encountered in
the literature and in teaching at international workshops for
advanced users of the clinical research literature: 1) key
concepts to reduce residual confounding in randomised trials;
2) methodological aspects regarding the choice and use of
primary and secondary outcomes; 3) sample size calculations
and the importance of event rates for these calculations; and 4)
how to present and use outcome information in randomised
trials.

KEY CONCEPTS TO REDUCE RESIDUAL CONFUNDING
IN RANDOMISED TRIALS
Highly valid trials provide estimates about treatment effects
that are as close as possible to the truth. These effect estimates
may be expressed as relative estimates (relative risks or odds
ratios) or as absolute effects (including natural frequencies, risk
differences or numbers needed to treat). There are a number of
methodological factors that can bias results, such as lack of
blinding of investigators who assess outcomes (e.g. when
assessing 6-min walk distance) or an inappropriate per-
protocol instead of an intention-to-treat analysis. These latter
factors are well described in other texts [5].

Another fundamental concern that may lead to systematic over
or underestimation of treatment effects is the presence of
confounding [6]. Confounding refers to an observed effect (e.g.
on mortality) that is not only due to the exposure of interest
(e.g. treatment with a long-acting bronchodilator) but also due
to some other outcome-determining factor (e.g. disease
severity). In “classic” observational epidemiology, a confoun-
der is a factor that is associated with the exposure (causally or
noncausally) and influences the outcome (e.g. an association
between yellow fingers from nicotine exposure and death is
completely confounded by smoking, which causes both yellow
fingers and death). An example of partial or incomplete
confounding is the influence of disease severity in observa-
tional studies on the association between bronchodilators and
death. Severely ill patients are more likely to receive a
bronchodilator and are also at a greater risk of death. In the
context of randomised trials, unbalanced groups (e.g. one
group suffers, on average, from more severe disease) can cause
confounding. Although it may be argued that this is not a
typical form of confounding but a form of random error
resulting from randomisation, the consequences remain
analogous: disease severity influences and, thereby, confounds
the observed effects. Unbalanced groups in randomised trials
are, therefore, not a form of random error that would reduce
precision of effect estimates but a source of residual confound-
ing that needs to be controlled for. It should be noted,
however, that a factor such as disease severity may also be of
interest because it could alter the effects of a treatment. The
latter, called effect modification, is particularly important for
clinicians because it may inform research for targeting
treatments to patient groups that benefit most. A prerequisite
for evaluating effect modification is, however, that groups are
evenly balanced for the effect modifier of interest (e.g. disease
severity) in order to allow for appropriate subgroup analyses
and interpretation according to well-defined criteria [7].

While there is no protection against imbalance of unknown
confounders, investigators can protect their study results
against known factors that may cause residual confounding.
We will describe three common approaches to control for
known and measurable confounders: restriction, pre-stratifi-
cation and statistical adjustment.

Restriction
Restriction is a simple method that describes the exclusion of
patients with characteristics that may introduce confounding
(e.g. patients with more severe disease), and is a very powerful
method of reducing the risk of confounding. However, it
comes at the cost of recruiting selected study populations,
which, in turn, may limit the applicability of findings.
Applicability should be distinguished from internal validity,
i.e. whether the study results in themselves are valid for the
setting, population, intervention and outcomes of the study
based on its design and execution. Applicability refers to
external validity and whether the results are valid when
applied outside of the trial environment. While a number of
approaches exist to evaluate applicability [8, 9], the GRADE
(grades of recommendation assessment, development and
evaluation) working group has provided a general conceptual
approach for the distinction between internal validity, or risk
of bias for a given body of evidence, and how directly the
findings of this evidence can be applied to the question in
healthcare [10, 11]. The concept of directness includes making
judgments about whether similar results could be expected
when transferring evidence from the trial PICO (population,
intervention, the comparison intervention (e.g. placebo or
alternative management) and outcomes) to the PICO of the
actual healthcare questions that requires an action. For
example, if a trial or a body of trials included only patients
with moderate COPD, are the results applicable to patients
with severe COPD (P); are results obtained with one inhaled
steroid expected to differ across the entire class and for
different doses (I); are the comparator interventions or
alternatives used in the trial the same in clinical practice (C);
and are the outcomes directly relevant to patients or is there
uncertainty about how important they are for decision making
(O). ROTHWELL [7] provides helpful specific examples that fall
into the PICO categories, such as genetic factors and
comorbidities, as examples for making judgments about the
directness of the population. Restriction will limit directness,
but if indirectness is judged to be of little concern or if the focus
is efficacy, it is a powerful means to reduce potential spurious
findings.

Pre-stratification
Pre-stratification is commonly used and means that random-
isation is, for example, performed separately for patients with
moderate and severe COPD. In this way, equal number of
patients with moderate and severe disease will be allocated to
treatment groups, reducing the risk of imbalanced groups.

Statistical adjustment
If groups are not well balanced for potential confounders,
analysts can still adjust for imbalances in the analyses. For
example, in a randomised trial that assessed the effects of respiratory rehabilitation on mortality, the control group had, on average, more severe disease, as expressed by a lower exercise capacity and poorer lung function, two factors known to be associated with mortality [12–14]. The effects of respiratory rehabilitation effects may be exaggerated because the average prognoses of the groups were not the same. Introducing and adjusting for measures of exercise capacity and lung function in the statistical analysis (for example in multivariate regression analyses) could account for these differences. However, it should be noted that statistical adjustment cannot eliminate the problem of unknown or unmeasurable confounders.

Restriction, pre-stratification and statistical adjustment are rarely used in COPD trials [1, 2]. For their optimal use, it is important to identify potential confounders during the design stage of a trial. Therefore, it is useful to explicitly describe the main relationship (fig. 1a) defined by the treatments and the primary outcome. All potential confounders, often suggested by prior research, should be listed and evaluated according to their potential to bias the primary outcome. Thereby, investigators can prioritise the most important confounders and decide how to control for them in the design phase. Control for confounding during the planning stage using randomisation, restriction and pre-stratification, and statistical adjustment during analysis of randomised trials, will reduce the type of confounding we have described.

**METHODODOLOGICAL ASPECTS REGARDING THE CHOICE OF PRIMARY AND SECONDARY OUTCOMES**

The CONSORT (Consolidated Standards of Reporting Clinical Trials) statement, a blueprint for the reporting of clinical trials, asks authors to identify a single primary outcome measure and a clear distinction from secondary outcomes [5]. The rationale is that investigators should specify which outcome they considered to be of greatest importance to guide interpretation of the trial. Specifying only one primary outcome prevents problems of interpretation associated with multiplicity of analyses. A second common and related reason to define a primary outcome relates to sample size calculation. The underlying reason is that confounders may differ by outcome. Consider figure 1a, which shows a trial investigating the effects of long-acting bronchodilators on mortality. Plausible confounders to control for include forced expiratory volume in 1 s (FEV₁; or, alternatively, the Global Initiative for Chronic Obstructive Lung Disease stage), previous exacerbations and exercise capacity as risk factors for mortality. In contrast, if health-related quality of life (HRQL) is of primary interest (fig. 1b), the most important confounders one might want to control for may include dyspnoea at baseline, previous exacerbations and comorbidities. Restriction and pre-stratification for the latter trial may differ substantially from the former. This simple example shows that the confounders that need to be controlled for by using the aforementioned methods depend on the primary outcome. As a consequence, trial results for secondary outcomes might be prone to residual confounding, particularly if trials are small.

**How to choose the primary outcome**

The most important criterion for choosing the primary outcome in trials should be the degree of importance of the outcome to patients. A number of methods exist for determining the importance or value of outcomes from the utility literature, but there is a general lack of agreement on the best method [11, 16]. However, integrative measures such as overall HRQL, exacerbations and mortality are logical primary outcomes in COPD trials. HRQL can be measured with good validity and reliability [17]. Furthermore, many validated HRQL instruments are also responsive to change [18]. However, in some instances investigators might want to gain more insights into pathophysiological processes and decide on primary outcomes that are not important for patients, but help in explaining the occurrence, mechanism or progression of disease.

**FIGURE 1.** Defining the main relationships between treatment and outcome. GOLD: Global Initiative for Chronic Obstructive Lung Disease.
Once investigators have selected a primary outcome, measurement considerations play an important role. Exacerbations, for example, can have a symptom-based [19] or an event-based definition [20]. Adjudication committees should decide, based on complete and unbiased data collection, whether patients experienced an exacerbation or not [21]. For the assessment of HRQL, instruments should be available and validated in the respective language. Finally, it is important to choose instruments for which guidance about interpretation of patient importance exists, such as thresholds for the minimal important difference (MID) [22]. The MID can also guide sample size calculations and trial interpretation [16].

Secondary outcomes
Secondary outcomes remain important because they may offer additional insights and offer hypothesis-generating results. In the TORCH (Towards a Revolution in COPD Health) study [21], for example, the investigators could confirm concerns that inhaled corticosteroids increase the risk of pneumonia. Thus, the use of secondary outcomes is critical for balancing benefits and downsides of treatments. While single trials can often not provide sufficient data on such secondary outcomes, they can then be summarised in well-performed systematic reviews that should be regularly updated to integrate new data. An example of the value of collecting data on such secondary outcomes is supported by the results of the UPLIFT (Understanding Potential Long-term Impacts on Function with Tiotropium) study [23], which inform the concerns raised in a recent systematic review and a larger registry study suggesting an increased risk of cardiovascular adverse effects and death from anticholinergic metered-dose inhalers in COPD [24, 25].

The number of patients and the deaths as secondary outcome in long-term tiotropium treatment trials increased from 3,247 and 97 in the systematic review by Singh et al. [24], to 8,294 and 1,038 after UPLIFT, respectively. The detailed reporting of the secondary outcome data in the UPLIFT study, thus, helped to inform the debate about the increased death rate by providing a large number of additional events.

However, results from secondary outcomes should be interpreted with caution because of several potential caveats, such as less careful ascertainment of the data. For example, mortality was a secondary outcome in the recent INSPIRE (Investigating New Standards for Prophylaxis in Reducing Exacerbations) trial that compared combined treatment (inhaled steroid plus long-acting ß-agonist) with a long-acting anticholinergic alone [26]. Although this trial showed very similar results for the primary outcome (exacerbation rates for both treatment regimens around 1.3 exacerbations per person-yr with p=0.66 for between-group difference) mortality was significantly lower in patients with combined treatment (3% versus 6% with long-acting anticholinergic; p=0.03). In the context of no difference in exacerbation rates and evidence from earlier trials it does not seem plausible that combined treatment lowers mortality compared to long-acting bronchodilator alone [27]. Indeed, a closer look at how death was ascertained in the INSPIRE trials reveals that death was only counted for patients who did not withdraw from treatment (65.5% with combined treatment and 58.3% with long-acting anticholinergic). Thus, it appears that one-third of patients did not enter the mortality analysis. Following the intention-to-treat principle protects against the bias that this per-protocol analysis may have caused [28, 29]. This example indicates that less stringent measurement of secondary outcomes may cause severe problems with missing data and render intention-to-treat analyses difficult.

In addition, the use of multiple secondary outcomes augments problems with interpretation and reporting. For example, how should we interpret contradicting results from secondary outcomes? Often, the number of measured secondary outcomes is so large that positive results emerge by chance. Methodological and statistical safeguards and solutions are available to address these concerns, but avoiding selective reporting of secondary outcomes will be the most important safeguard. Secondary outcomes should be selected to complement and enhance the plausibility of the main aim of the trial and inform decision making. In the TORCH study [21], for example, exacerbations and FEV1 were secondary outcomes that are associated with death in prognostic studies. The effect of combination treatment was consistent across these outcomes. One might argue that the reduction in mortality compared with placebo, although it just failed to reach conventional statistical significance, can be explained by a reduction in exacerbations and a slower decline in lung function, both of which are associated with lower mortality and suggest disease modification.

Results based on secondary outcomes require careful evaluation before they should inform clinical practice. The reasons include: that data collection might be less stringent and robust than for the primary outcome; that selective reporting of secondary outcomes showing significant results is frequent; and that methodological problems, such as insufficient control for confounding and the use of per-protocol analyses, can limit the validity of results from secondary outcomes. Finally, systematic reviews including those of secondary outcomes should become a standard approach to systematically summarising findings across studies.

SAMPLE SIZE CALCULATIONS AND THE IMPORTANCE OF EVENT RATES
Once investigators have determined the primary outcome and how to control for residual confounding, the sample size should be estimated to ensure adequate precision of the effect estimates. The TORCH study selected mortality as the primary outcome [21]. The investigators determined the required sample size to detect an expected absolute difference in mortality after 3 yrs between the placebo group (17%) and the group with combined treatment (12.7%) with 90% power. The observed mortality rates were 15.2% (placebo group) and 12.6% (combined treatment group), respectively. Thus, the assumed difference of 4.3% (equal to 17% minus 12.7%, for a relative risk reduction of 25.3%) for the absolute reduction in death was much larger than observed. Analogously, the assumed relative risk reduction was much larger than the actually observed relative effect (15.2% minus 12.6% equals 2.6%; 2.6% divided by 15.2% equals an observed effect of 17.1%).

Unfortunately, overestimation of both event rates and relative treatment effects is common for sample size calculation of many trials. Definition of what constitutes an important risk
reduction is challenging for binary outcomes such as mortality. In practice, a steering committee decides what constitutes a relevant treatment effect, but this is often not based on solid data. While there is no simple solution for this dilemma, the estimates should be realistic and relative treatment effects of $>20\%$ are uncommon and should, therefore, be applied with caution [30].

Furthermore, beyond sample size, one of the main factors determining sample size calculations are event rates; that is, the number of patients who experience an outcome. The following calculation exemplifies this problem: consider a large randomised trial of 20,000 patients randomised into two groups, treatment and placebo, of 10,000 patients each. One would believe that trials of this magnitude should be able to detect large effects, such as relative risk reductions of 50\%, with great certainty. However, if the event rate is low (e.g. one event in the treatment group and two events in the placebo group, as an extreme example shown in table 1), the results would not be statistically significant ($p=0.56$). There would be great uncertainty about the true effect, making a very large reduction in risk of 95\% equally likely as a relative risk increase of 450\%.

Conversely, a much smaller trial with 2,000 patients (1,000 in the treatment and the placebo group each) and 100 and 200 events, respectively, will yield the same relative risk of 0.5 (p<0.001) with sufficiently precise confidence intervals of 0.37–0.60 to indicate that a large effect exists (table 2).

In a more realistic example, investigators might want to observe a relative risk reduction in exacerbation rates of 20\% over a treatment period of 1 yr; assuming a baseline event rate of 20\% in the control group and a 16\% event rate in the treatment group, around 3,000 patients are needed (80\% power). If the baseline event rate is 40\% only around 1,200 patients are needed to show a statistically significant relative risk reduction of 20\%. Exacerbation baseline rates are determined by inclusion and exclusion criteria, the definition of exacerbations and length of follow-up. Investigators should reflect carefully on these three parameters during the planning stage, as they determine sample size, funding requirements and feasibility. For continuous outcomes, such as HRQL data, defining both important treatment effects and sample size is simpler when the MID has been established or other thresholds for interpretability exist [22].

Multi-arm trials such as TORCH offer an additional challenge. The single drug groups (bronchodilator and inhaled corticosteroid) were apparently not considered for sample size calculations. The four-armed TORCH study opens, thus, the debate whether sample size calculations should be based on the most important comparison or on more than one comparison. For clinicians, the relevant question would have been whether to treat patients with a single drug (bronchodilator) or with a combination (bronchodilator plus inhaled steroid) rather than the comparison between combination treatment and placebo. The focus on the latter comparison may have been guided by the believe that trials should always include a “least possible treatment” arm, but investigators and those using the literature should realise that the placebo control may be necessary only for add-on treatments. For example, the TORCH trial could have been powered to show that combination treatment is superior to single bronchodilator treatment if the single treatment arm had also received a placebo. The trial could also have been powered to explore whether single bronchodilator treatment is not inferior to combination treatment.

In superiority trials that focus on the superiority of a drug over placebo, the power to detect expected differences is commonly set at 80\%. In order to yield more precise results, the TORCH trial set the power at 90\% [21], which increases sample sizes

| TABLE 1 | The relationship of event rates to relative and absolute risk reductions |
|-----------------|-------------------------------|-------------------|
| **Patients with outcome n** | **Patients without outcome n** | **Event rate (risk)** |
| Treatment group | 1 | 9999 | 0.0001 |
| Placebo group | 2 | 9998 | 0.0002 |
| Absolute risk reduction or risk difference | 0.0001 (95% CI -0.0002–0.0004) |
| Relative risk | 0.50 (95% CI 0.05–5.51) |
| Relative risk reduction | 50% (95% CI -95%–450%) |

| TABLE 2 | Smaller trials with high event rates can provide narrow confidence intervals |
|-----------------|-------------------------------|-------------------|
| **Patients with outcome n** | **Patients without outcome n** | **Event rate (risk)** |
| Treatment group | 100 | 900 | 0.1 |
| Placebo group | 200 | 800 | 0.2 |
| Absolute risk reduction or risk difference | 0.1 (95% CI 0.07–0.13) |
| Relative risk | 0.50 (95% CI 0.40–0.63) |
| Relative risk reduction | 50% (95% CI 37%–60%) |
substantially. In noninferiority or equivalence trials, however, investigators are less flexible in their choice of power. The power should be set at ≥ 90% in order to minimise the problem of wrongly assuming equivalence if the effects of two interventions actually differ (commonly referred to as type II or β error) [16].

Finally, in the realistic situation of limited funding, sample size calculations will depend on issues of feasibility, and many approaches to sample size calculations exist [30]. We described that fundamental requirements for sample size calculations include: careful selection of the primary outcome; event rates; consideration of the main comparison, in the case of multi-arm trials; and the assumption of effect sizes based on the MID or on realistic estimates of relative or absolute risk reductions.

HOW TO PRESENT AND USE INFORMATION OF BENEFITS AND OUTCOMES IN CLINICAL TRIALS

COPD trials such as TORCH raise, as do other multi-arm studies, several issues for reporting and interpretation. The investigators drew their main conclusions on the basis of comparisons between combined treatment and placebo. Combined treatment appears superior to placebo. However, how should we interpret the comparison between combined and single bronchodilator treatment, which may indeed be the more important clinical question? There were no statistically significant differences between these two treatment options with respect to mortality and severe exacerbations (hospital admissions), but combined treatment significantly reduced any (moderate and severe) exacerbations and moderate exacerbations and reduced the decline of FEV1. Differences in HRQL were statistically significant but these differences were well below the MID; only a comparison of the proportions of patients achieving the MID would inform the discussion. Finally, combined treatment significantly increased the risk of pneumonia (19.6% over 3 yrs) compared with single bronchodilator treatment (13.3%). Therefore, one could argue that single bronchodilator treatment should be favoured because there was no important difference in the primary outcome. Furthermore, there were no differences in other outcomes important for patients, such as hospital admissions or HRQL, while the safety profile was more favourable for single bronchodilator treatment.

This example illustrates that informative but complex trials such as TORCH need to be viewed by balancing all important outcomes, in particular in the absence of a clearly dominating mortality difference. However, how can we ensure that clinicians and patients are informed about all results? The foregoing question raises the more important questions of how should physicians and patients be informed and how should they balance the benefits and downsides of single bronchodilator and combined treatment? We emphasised that a prerequisite for balanced decisions is that reporting is as complete as it is in the case of the TORCH trial. Interpretation should include all comparisons and all outcomes [5]. This would enable investigators to develop decision aids, where, based on the entire body of evidence from all available trials and summarised in high-quality systematic reviews, the benefits and downsides are illustrated comprehensively [31]. Decision aids are particularly valuable for value-sensitive decisions in which the balance of benefits and downsides is not straightforward and patients have different views on outcomes and treatments. Whether patients benefit from such an informed decision making requires, however, testing in additional trials.

CONCLUSION

The design and conduct of COPD trials requires a clear definition of the main relationship between treatment and primary outcomes, identification of confounders and selection of means to optimally control for these confounders during the design phase. Selection of the primary outcome should be guided by the importance to patients. Realistic estimates of treatment effects should inform trials. Secondary outcomes will provide complementary information to explain or describe consequences of the effects observed for the primary outcome, but should be interpreted with caution because they often provide limited data or are incompletely reported. Finally, essential steps to transfer the results from research to practice requires complete reporting of results and developing tools such as decision aids that convey benefits and downsides of treatments in an informative way to patients and clinicians, in order to support their shared decision making.

SUPPORT STATEMENT

M.A. Puhan is supported by a career award of the Swiss National Science Foundation (3233B0/115216/1). H.J. Schünemann is funded by a European Commission grant: the human factor, mobility and Marie Curie actions (Scientist Reintegration Grant; IGR 42192).

STATEMENT OF INTEREST

Statements of interest for M.A. Puhan and H.J. Schünemann can be found at www.erj.ersjournals.com/misc/statements.dtl

ACKNOWLEDGEMENTS

The authors are grateful to L. Fabbri (University of Modena and Reggio Emilia, Modena, Italy) for suggesting this topic and reviewing an earlier version of the document.

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