Bias and Loss to Follow-Up in Cardiovascular Randomized Trials: A Systematic Review

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BACKGROUND: Loss to follow-up (LTFU) is common in randomized controlled trials. However, its potential impact on primary outcomes from cardiovascular randomized controlled trials is not known.

METHODS AND RESULTS: We conducted a prospective systematic review (PROSPERO: CRD42019121959) for randomized controlled trials published in 8 leading journals over 5 years from January 2014 to December 2018. Extent, reporting, and handling of LTFU data were recorded, and the proportion of a trial's primary outcome results that lose statistical significance was calculated after making plausible assumptions for the intervention and control arms. These assumptions could drive differential treatment effects between the groups considering relative event incidence between LTFU participants and those included in the primary outcome. We identified 117 randomized controlled trials of which 91 (78%) trials reported LTFU, 23 (20%) reported no LTFU, and 3 (3%) trials did not report on whether LTFU occurred. The median percentage of study participants lost to follow-up was 2% (interquartile range, 0.33%–5.3%). Only 10 trials (9%) had a low cluster of risk factors for impairment in trial quality. The percentage of trials losing statistical significance varied from 2% when the relative event incidence for LTFU between the randomized groups was 1 for the intervention arm and 1.5 for the control arm to 16% when the relative event incidence was 3 for the intervention arm and 1 for the control arm.

CONCLUSIONS: Almost 1 in 6 (16%) cardiovascular randomized trials published in leading journals may have a change in the primary outcome if plausible assumptions are made about differential event rates of participants lost to follow up. There is scope for improvement arising from LTFU in randomized trials in cardiovascular medicine.

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Key Words: bias ■ loss to follow-up ■ outcome ■ outcome and process assessment ■ patient dropout ■ randomized controlled trials ■ relative risk

The gold standard assessment of a medical intervention involves assessment in a randomized controlled trial (RCT). Randomization balances the distribution of any known or unknown potential confounding factors between treatment arms. This mitigates the possibility of selection bias, especially if the participants’ group allocations are concealed. Blinding of patients, therapists, and outcome assessors is an additional useful tool to prevent bias. Open-label clinical trials are often unavoidable if blinding of patients and therapists is not possible. Clinical guidelines may be influenced by biased clinical evidence leading to undesirable impacts on patients, healthcare providers, and funders. Up to 80% of contemporary clinical RCTs fail to achieve complete follow-up. This important factor may affect the integrity of study conclusions. If participants are lost and the characteristics of such
participants associate with clinical events, then bias can arise. This is particularly relevant in open-label studies in which assessors know the group allocations of the participants. Loss to follow-up (LTFU) in this scenario could favor the intervention arm and neutralize the benefit of randomization. It is plausible that attrition bias associated with LTFU drives either overestimation or underestimation of treatment effects.

Classification of LTFU and recommendations for dealing with LTFU have been made. Crucially, however, the contemporary prevalence and effects of LTFU within cardiovascular trials is not known. This prospective systematic review and meta-analysis was designed to analyze the prevalence and potential impact of LTFU in cardiovascular RCTs. The primary aim was to assess the proportion of trials in which the primary efficacy end point would change if plausible assumptions were made about participants who were unaccounted for in the original analysis. In addition, we assessed estimates of treatment effect according to the extent, reporting, and handling of LTFU and trial characteristics associated with LTFU.

METHODS

Eligibility

All supporting data are available within the article and its online supplementary files. Ethics approval was not required. We predefined reports as being eligible for inclusion in this analysis if an RCT in cardiovascular disease was described and published in one of the 5 leading general medical journals and 3 cardiology journals with the highest impact factors (Annals of Internal Medicine, BMJ, JAMA, Lancet, New England Journal of Medicine, Circulation, European Heart Journal, and Journal of the American College of Cardiology). A 5-year publication period was set from 2014 to 2018. An additional inclusion criterion was if a patient-important binary primary outcome was statistically significant at a 2-sided α of 0.05. The rationale behind focusing on statistically significant trials in major journals only is that the results of these trials are most likely to influence clinical guidelines. Therefore, a change in significance of a risk ratio due to bias might affect patient care to an important extent. Cluster trials, crossover trials, N-of-1 trials, and trials reported in research letters were excluded. Equivalence and noninferiority studies were excluded unless the authors prespecified testing for superiority. Reports describing secondary analyses of randomized trials were excluded.

A patient-important outcome was defined as an outcome that would be undesirable for a patient to experience and the patient would thus try to prevent it by undergoing an effective treatment. Mortality and morbidity are examples of outcomes that were included. Surrogate outcomes were considered as nonpatient important (Data S1). The protocol was registered on PROSPERO (CRD42019121959).

Literature Search

Reports of RCTs were identified from Medline and Embase using OVID (Data S2). The search was restricted to clinical RCTs in cardiovascular disease published in the selected journals between 2014 and 2018. Trials were considered statistically significant if the 2-sided 95% CI of an estimate of the relative risk did not include 1.0 or if the 2-sided P value for superiority was <0.05 when no CI was reported. A calibration exercise was performed before the search. One reviewer identified and reviewed the potentially eligible reports based on an agreed screening form (Data S3). The list of included and excluded reports was provided to the 2-person reviewer team after screening. Disagreements were resolved by consensus, with the assistance of a third reviewer as required.
Data Collection
Data were extracted based on an agreed data extraction form (Data S4). The primary outcome selected for the review was the one specified within the report. If the report specified both significant primary efficacy and safety outcomes, the primary efficacy outcome was selected. If multiple primary outcomes were specified, the statistically significant outcome in the highest category on the outcome hierarchy was selected (Data S1). If both intention-to-treat and per-protocol analyses were reported, we considered the statistical significance of the former; if both unadjusted and adjusted analyses were reported, the statistical significance of the former was considered. Data on study background, general characteristics, methodological quality, the extent of LTFU, its reporting, and its handling in the analysis related to the primary outcome were extracted. Patients were considered as LTFU if they were mistakenly randomized with inappropriate postrandomization exclusion; did not receive the intervention or adhere to treatment, with inappropriate postrandomization exclusion; withdrew consent; crossed over to another arm but were not included in the analysis; or lost contact. Trials were categorized by subspecialty focus: electrophysiology, heart failure, interventional cardiology, cardiac surgery, general cardiology, and cardiovascular imaging.

Statistical Analysis
The analysis is explained in more detail in the online supplement (Data S5). Methodological and reporting quality of the included trials was assessed, as suggested by Bikdeli et al14 and the Cochrane risk-of-bias assessment tool.15 The extent of LTFU was calculated as the percentage of LTFU in each trial from each arm (intervention and control). The ratio of LTFU rate to primary event rate was also reported. A univariable random-effects metaregression analysis was conducted using the log odds of participants lost to follow-up as the dependent variable and general trial characteristics and methodological characteristics as independent variables.

The potential impact of LTFU on the primary outcome analysis was evaluated by making assumptions about the outcomes in LTFU participants (Data S6). An estimation algorithm proposed by Akl et al17 was adopted with relative incidence of outcomes in LTFU patients compared with patients who were followed-up (RILTFU/FU), ranging from 1 to 3. In addition, the following common assumptions were used for calculations: none of the participants lost to follow-up had the event; all participants lost to follow-up had the event; none of those lost to follow-up in the treatment group had the event, and all those lost to follow-up in the control group did (best case scenario); all participants lost to follow-up in the treatment group had the event and none of those in the control group did (worst-case scenario).

For each trial, 2x2 tables were constructed for the collected data for the calculation of risk ratios associated with each assumption. The percentage of trials with their primary outcome becoming nonsignificant was calculated based on the assumptions and definition of statistical significance reported above. Trials with no LTFU were excluded in the primary analysis but included in a sensitivity analysis. An additional prespecified sensitivity analysis stratified by type of intervention was conducted. Paired differences in

Figure 1. Search and screening approach.
Flow of trial reports identified and screened in this analysis is shown. The search recovered 3668 reports; 1873 reports were screened after removing duplicates; 117 reports were included after screening, and reasons for exclusion are stated in text. FLUKE indicates Follow Up Loss Effect Upon Skewing Evidence.
proportions between interventional cardiology trials and those of other cardiology subspecialties were also assessed based on different assumptions.

Table 2. Methodological and Reporting Quality Assessment of the Included Trials

| Factors                                      | Trials at Risk of Bias (n=117), No. (%) |
|----------------------------------------------|----------------------------------------|
| inadequate allocation sequence concealment  | 63 (54)                                |
| no blinding of patients†                     | 78 (65)                                |
| early stop                                   | 9 (8)                                  |
| not using intention-to-treat analysis†       | 29 (25)                                |
| absence of protocol†                         | 31 (26)                                |
| without explicit statement about status of LTFU | 43 (36)                               |

LTFU indicates loss to follow-up.

Allocation concealment defined as to the person enrolling participants does not know in advance which treatment the next person will get which usually involves the use of computer algorithms. It seeks to prevent selection bias by protecting the assignment sequence until allocation, and can always be successfully implemented. It is considered to be adequate according to the definition reported by Jüni et al.Allocation defined as to the withholding information about the assigned interventions from people involved in the trial who may potentially be influenced by this knowledge; blinding is performed to prevent performance and ascertainment bias by protecting the sequence after allocation and cannot always be implemented. It is considered to be adequate only if clearly indicated.

Intention to treat analysis defined as an analysis that included all randomized participants in the analysis who are all retained in the group to which they were allocated.

Consider as absence if the protocol is not published before or is included as appendix beside the main report.

Among the trials that LTFU occurred (n=91)†:

- Separately reported in 2 arms: 70 (77)
- Compared the LTFU group baseline characteristics with not LTFU: 0 (0)
- Implication of LTFU discussed: 6 (7)

Distribution of trials according to methodological and reporting quality assessments that might impair the outcomes of the trial.

Distribution of trials according to the number of methodological and reporting quality characteristics (methodological confounders) they possess after the assessment: 9% of the trials had none of the methodological confounders (n=10), 21% of the trials possessed 1 methodological confounder (n=24), 32% of the trials possessed 2 major methodological confounders (n=38), and 26% of the trials possessed 3 major methodological confounders (n=30). In addition, 12% of the trials had >3 methodological confounders. This list of methodological confounders analyzed included the following: (1) inadequate allocation sequence concealment, (2) no blinding of patients, (3) early stop of trial, (4) not using intention-to-treat analysis, (5) absence of protocol, and (6) no explicit statement about status of loss to follow-up.
RESULTS

After excluding duplicates and screening for eligibility, 117 studies were included from a total of 3668 from the initial search (Figure 1). The list of the 117 studies included in this analysis is provided in Table S1.18–134 The mean age of 407,229 study participants was 64.2 years (30% female). The trial subspecialties were electrophysiology (19%), heart failure (3%), interventional cardiology (28%), cardiac surgery (3%), general cardiology (44%), and cardiovascular imaging (3%). Baseline study characteristics of the included trials are reported in Table 1.

Assessment of the Methodological Quality of the Trials

The analytical methods that were used for handling LTFU in the primary analysis of the included trials are presented in Table 1. The most commonly used method
was censoring at time of LTFU in a time-to-event analysis (N=75; 64%). Two trials (2%) assumed that no LTFU participants experienced events, whereas 10 (8%) used complete case analysis and 2 (2%) used a worst-case scenario in which only the control arm had events. Two trials (2%) used multiple imputation, whereas none reported using inverse probability weighting.

Regarding the reporting of LTFU, 85 (73%) used a Consolidated Standards of Reporting Trials (CONSORT) diagram. Seventy (77%) trials reported...
that LTFU occurred in the intervention and control arms separately. However, none of the trials compared baseline characteristics of LTFU participants with followed-up participants. The implications of LTFU are discussed in 6 trials (7%).

Table 2 and Figure 2 demonstrated the number of trials meeting the characteristics (methodological confounders) for impairment in the quality of trial design. Allocation concealment was adequate in 54 trials (46%). Patients were blinded adequately in 41 trials (35%). In 9 trials (8%), enrollment was discontinued prematurely. Twenty-nine trials used an intention-to-treat analysis (25%). Thirty-one trials (26%) provided a protocol. Forty-three trials (36%) did not state the status of LTFU explicitly in the report. Only 10 trials (9%) were free from any methodological confounders that might impair the methodological quality.

Random-effects metaregression analysis (Table 3) suggested that inadequate or unclear concealment of allocation was associated with an increase in odds of LTFU (odds ratio, 2.37 [95% CI, 1.42–3.97]; \( P = 0.001 \)). Increasing sample size (\( P = 0.039 \)) and duration of follow-up (\( P = 0.002 \)) also increased the odds of LTFU. Finally, the odds of LTFU was decreased in nonsurgical or noninterventional trials (odds ratio, 0.50 [95% CI, 0.30–0.85]; \( P = 0.01 \)).

Table 4. Percentage of 91 Trials in Which Results Would Lose Significance Under Different Assumptions on the Outcomes of LTFU Participants in Intervention and Control Arms

| N=91 | R_{LTFU/FU} | 3 | 2 | 1.5 | 1 |
|-------|-------------|---|---|----|---|
| RI_{LTFU/FU} (Intervention) | 1 | 3 | 3 | 2 | 4 |
| RI_{LTFU/FU} (Control) | 1.5 | 3 | 2 | 3 | 4 |
| RI_{LTFU/FU} (Intervention) | 2 | 4 | 3 | 4 | 12 |
| RI_{LTFU/FU} (Control) | 3 | 3 | 9 | 10 | 16 |

Among the 91 trials, percentages of results that would lose significance under less plausible assumptions: (1) none of the LTFU participants had the event, 4%; (2) all the LTFU participants had the event, 11%; (3) none of those lost to follow-up in the treatment group had the event, and all those lost to follow-up in the control group did (best case scenario), 3%; (4) all participants lost to follow-up in the treatment group had the event, and none of those in the control group did (worst case scenario), 33%. FU indicates follow-up; LTFU, loss to follow-up.

\(*_{RI_{LTFU/FU}} is the relative event incidence among those with LTFU compared with those followed up.

Figure 5. Bias and loss to follow-up in randomized controlled trials in cardiovascular medicine. Assumptions being made toward the outcome of LTFU in each trial from the search and the subsequent calculation made. In total, 117 trials from 8 journals covering 407,229 patients from 2014 to 2018 were recovered. Assume participants were randomized to intervention and control, respectively; 3 had events from each arm and 3 dropouts from each arm. From the figure, dotted transparent figures denote LTFU participants, whereas red dotted figures denote LTFU participants being assumed with event. The plausible assumptions being made toward the LTFU was based on relative event incidence and a formula detailed in Data S6. The number of events were assumed based on the reported formula with incidence ranging from 1 to 3. Calculations of how many trials’ relative risks lost significance after making the assumptions were run subsequently. Ann of Intern Med, Annals of Internal Medicine; Eur Heart J, European Heart Journal; JACC, Journal of the American College of Cardiology; LTFU, loss to follow-up; NEJM, New England Journal of Medicine.
Extent of Loss to Follow-Up
Among the 117 included trials, 91 (78%) reported LTFU. Twenty-three trials reported no LTFU (20%), and 3 trials did not report whether there was LTFU (3%). Of the trials with LTFU, the median percentage of LTFU was 2% (interquartile range [IQR], 0.3%–4.8%) in the intervention arm, 1.99% (IQR, 0.3%–5.4%) from the control arm, and 1.96% (IQR, 0.33%–5.3%) overall. The median difference between the intervention and the control groups was not significant ($P=0.978$; Figure 3). The medians for the ratios of LTFU to events were 0.12 (IQR, 0.03–0.33) in the intervention arm, 0.11 (IQR, 0.02–0.42) in the control arm, and 0.11 (IQR, 0.03–0.41) overall. A value of 0.12 means that $\approx 1$ participant is LTFU when every 10 participants experience the primary outcome. However, the difference between the ratio of the intervention and the control groups was not significant ($P=0.473$; Figure 4).

Potential Impact of LTFU

Percentage of Trials Losing Significance
For the 4 common assumptions in which all 91 trials were included, the percentages of trials that lost significance were 4% (no participants lost to follow-up had the event), 11% (all participants lost to follow-up had the event), 3% (best-case scenario), and 33% (worst-case scenario).

Considering the relative event incidence analysis method, Table 4 shows the percentage of eligible trials that lost significance across a range of assumptions for the event incidence among intervention and control arms (Figure 5). The percentage varied from 2% to 16%. Figure 6 shows an inverse-proportion relationship of the trials losing significance with the percentage of LTFU under the best and worst assumptions made by the relative event incidence analysis method.

Results of the prespecified sensitivity analysis on the subspecialties are reported (interventional cardiology versus others) in the online Data Supplement. There was a significant difference in the proportion of trials losing significance between interventional cardiology and other subspecialties (difference, 4.35% [95% CI, 0.295%–8.41%]; $P=0.0369$; Figure S1 and Table S2).

DISCUSSION

We found considerable variation in the extent and reporting of LTFU data in contemporary cardiovascular clinical trials. We observed that certain characteristics of clinical trials—notably, inadequate or unclear allocation concealment, length of follow-up, sample size, and type of intervention—were associated with increased odds of LTFU. Importantly, the primary result

Table 5. Summary of the Important Common Issues for LTFU and Guidance in Conducting Trials and Reporting Trial Results

| Issues That Should Be Noted               | Guidance                                                                 |
|------------------------------------------|--------------------------------------------------------------------------|
| Inadequate or unclear allocation concealment | If allocation concealment forms part of the trial design, then effective approaches to achieve allocation concealment include using a matched placebo (visually identical to the active treatment); central randomization (performed at a site remote from the trial’s location); sequentially numbered, sealed, opaque envelopes |
| Large sample size and long follow up duration | LTFU increases with larger trial sample size, hence investigators should be aware and mitigate the number of LTFU for increase in sample size and 1-y increase in duration |
| Reporting of LTFU                        | Investigators should strive to reduce the number of LTFU. A CONSORT diagram should be included for readers. When LTFU has occurred, baseline characteristics, and extent of follow up duration before exclusion should be reported in the manuscript or supplement. The implications of LTFU should also be discussed in the manuscript. Time of dropout can be noted on a supplement or in the result paragraph or on the CONSORT diagram for readers to know the extent of follow up before dropout |

CONSORT indicates Consolidated Standards of Reporting Trials; LTFU, loss to follow-up.
in 1 of 6 trials might change if reasonable assumptions were made about the end point in patients with LTFU.

The inverse-proportion relationship noted in Figure 6 suggests that the impact of a small proportion of LTFU might be overlooked by investigators. More than one third of trials did not achieve effective blinding among either the participants or the site investigators. This finding is important because ineffective blinding is associated with overestimation of true treatment effects. Allocation concealment was inadequate or unclear in more than half of the trials. Conversely, an intention-to-treat analysis was used in 75% of trials, which minimizes the effects of attrition bias. Just over half of the trials included an explicit statement about LTFU, and >70% of the trials included a CONSORT flow diagram. Notably, baseline demographics on the LTFU, and >70% of the trials included a CONSORT half of the trials included an explicit statement about LTFU. We suggest that information on participants with LTFU should be included by authors in an appendix or in a defined column in a table of the trial participants’ characteristics (Table 5). Inverse probability weighting can be a helpful way of handling LTFU participants’ data, but it is not used in any of the included trials. Most trials did not impute data for LTFU participants. We noted a significant association between inadequate or unclear allocation concealment and increased odds of LTFU. This could be explained by less stringently implemented processes in trials with inadequate or unclear allocation concealment, including suboptimal measures for following up participants.**

**Strengths and Weaknesses of the Study**

Our study has several strengths. First, the forms for screening of the trials and related data collection were established before the start of the data collection process. In addition, the calibration exercise was completed upfront as a preparatory step intended to increase accuracy for the screening and data collection. Second, a range of assumptions was made for the participants with LTFU and explored the potential effect of LTFU on the estimate of the effect of the intervention, including whether or not the trial met statistical significance on its primary outcome and the change in the relative risk ratio and number of outcome events. The effect is focused on cardiovascular trials. Our analysis depends on the accuracy and clarity of the included reports. Generalizability is also an issue. We focused our analysis on 8 journals’ publications during a 5-year period (2014–2018). A wider inclusion strategy with more journals (with lower impact factors) and trials with a nonsignificant primary outcome result might have returned different results. Our findings might underestimate the true effect of LTFU in the effect estimate if a wider range of RCTs were included. Our review included trials with binary data only because of the design of the review analysis, which might further weaken the generalizability of the results. Time of dropout can be a factor influencing the LTFU effect because early dropouts can influence the result to a larger extent than late dropouts. However, exact time of dropout is not noted in the reports, and we are unable to stratify the effect.

**Implications**

Investigators and sponsors should strive to reduce the number of participants with LTFU. The higher the LTFU, the more uncertainty increases around the treatment effect estimate and the potential for a false result. In the unfortunate event that LTFU happened, its impact can be estimated using the worst assumption (Data S7). As for the reporting of LTFU, editors may consider requiring authors to provide a fully informative and transparent report on participant LTFU including the inclusion and exclusion criteria of patients, which is in line with CONSORT guidelines. Specifically, investigators should provide information on participants with LTFU including their baseline characteristics, reasons for LTFU, and duration of follow-up before exclusion and then compared with those who completed follow-up. This information could be published as an appendix. Implications of LTFU should be discussed when LTFU has occurred (Table 5). This review provides estimates of the probability that the primary analysis of cardiovascular trials could lose statistical significance when LTFU events are taken into account by making appropriate estimate of event incidence. Although the 4 less plausible but commonly used assumptions may not eventuate, they can be taken as the upper limit of change in trial significance. Early LTFU has a more influential effect on the analysis than late LTFU near the overall study duration, which highlighted the need for investigators to stratify LTFU by follow-up extent. Future studies can look at the extent of change in treatment effect in relation to the LTFU proportion and event number and the effect of partial and full LTFU defined as difference in the extent of follow-up before exclusion. The influence of dropout time on LTFU effect can be explored for assessing the possibility of systemic inclusion of patients accounting for early dropouts.

**CONCLUSIONS**

Almost 1 in 6 (16%) cardiovascular randomized trials published in leading journals may have a change in the primary outcome if plausible assumptions are made about differential event rates of participants lost to follow-up. There is scope for improvement arising from
LTFU in randomized trials in cardiovascular medicine. Bias minimization through mitigation of participants lost to follow-up offers the opportunity to enhance the value of randomized trials.

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Supplementary Materials
Datas S1–S7
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Data S1 - Hierarchy of outcomes relative to patient importance in FLUKE

I. Mortality
   a. All-cause mortality
   b. Disease specific mortality

II. Morbidity
   a. Cardiovascular major morbid events
   b. Other major morbid events (e.g. Revascularization)
   c. Onset/recurrence/relapse/remission of diseases
   d. Hospitalization, medical and surgical procedures
   e. Infections
Data S2- Search strategy for Medline and Embase using OVID interface

Medline
1 exp Myocardial Ischemia/
2 (MYOCARDS$4 adj4 (ISCHAEMIS$2 or ISCHEMIS$2)).tw.
3 exp Coronary Artery Bypass/
4 ((ISCHAEMIS$2 or ISCHEMIS$2) adj4 HEART).tw.
5 CORONARY.ti,ab.
6 exp Coronary Disease/
7 exp Myocardial Revascularization/
8 exp Myocardial Infarction/
9 (MYOCARDS$5 adj4 INFARCT$5).tw.
10 (HEART adj4 INFARCT$5).tw.
11 exp Angina Pectoris/
12 ANGINA.tw.
13 exp Heart Failure/
14 (HEART adj6 Failure).tw.
15 or/1-14
16 exp Heart Diseases/
17 (Heart adj4 disease$2).tw.
18 MYOCARDS$5.tw.
19 CARDIAC$2.tw.
20 CABG.tw.
21 PTCA.tw.
22 (STENT$4 and HEART).tw.
23 Heart Bypass, Left/ or Heart Bypass, Right/
24 CARDIOLOGY SERVICE, HOSPITAL/ or CARDIOLOGY/
25 or/16-24
26 15 or 25
27 Randomized controlled trial.pt.
28 randomized controlled trial/
29 (random$ or placebo$).ti,ab,sh.
30 ((singl$ or double$ or triple$ or treble$) and (blind$ or mask$)).tw,sh.
31 or/27-30
32 (retraction of publication or retracted publication).pt.
33 31 or 32
34 (ANIMALS not HUMANS).sh.
35 33 not 34
36 35 and 26
37 bmj.jn
38 “Annals of Internal Medicine”.jn.
39 jama.jn.
40 lancet.jn.
41 “new england journal of medicine”.jn.
42 36 and 37
43 36 and 38
44 36 and 39
45 36 and 40
46 36 and 41
47 european heart journal.jn.
48 circulation.jn.
49 journal of the American college of cardiology.jn.
50 36 and 47
51 36 and 48
52 36 and 49
53 42 or 43 or 44 or 45 or 46 or 50 or 51 or 52
54 limit 53 to yr="2014-2018"
EMBASE

1 Heart Disease/
2 (MYOCARDS$4 adj2 (ISCHAEMIS2 or ISCHEMIS2)).tw.
3 ((ISCHAEMIS2 or ISCHEMIS2) adj4 HEART).tw.
4 Coronary Artery Disease/
5 Transluminal Coronary Angioplasty/
6 (CORONARY adj4 (DISEASE$2 or BYPASS$2 or THROMBO$5 or ANGIOPLAST$2)).tw.
7 Heart Infarction/
8 (MYOCARDS$4 adj2 INFARCT$5).tw.
9 (HEART adj2 INFARCS5).tw.
10 Heart Muscle Revascularization/
11 Angina Pectoris/
12 ANGINA.tw.
13 (HEART adj2 FAILURE).tw.
14 (HEART adj2 DISEASE$2).tw.
15 CARDIACS$2.tw.
16 CABG.tw.
17 PTCA.tw.
18 (STENT$4 and HEART).ti,ab.
19 Extracorporeal Circulation/
20 cardiology/
21 or/1-20
22 Randomized Controlled Trial/
23 Single Blind Procedure/
24 Double Blind Procedure/
25 Crossover Procedure/
26 22 or 23 or 24 or 25
27 (random$ or factorial$ or crossover$ or placebo$ or (cross adj over) or assign$).ti,ab.
28 ((singl$ or double$ or triple$ or treble$) and (blind$ or mask$)).ti,ab.
29 controlled clinical trial*.ti,ab.
30 28 or 26 or 27 or 29
31 21 and 30
32 (animal$ not human$).sh,hw.
33 31 not 32
34 bmj.jn
35 “Annals of Internal Medicine”.jn.
36 jama.jn.
37 lancet.jn.
38 “new england journal of medicine”.jn.
39 european heart journal.jn.
40 circulation.jn.
41 journal of the American college of cardiology.jn.
42 33 and 34
43 33 and 35
44 33 and 36
45 33 and 37
46 33 and 38
47 33 and 39
48 33 and 40
49 33 and 41
50 or/42-46
51 or/47-49
52 limit 51 to article or article in press or conference paper
53 50 or 52
54 limit 53 to yr =”2014-2018”
Data S3 – FLUKE study data screening form

**FLUKE study: Data Screening Form**

| Screener initials: | Study ID: | Author, year: _______________ |
|--------------------|-----------|-----------------------------|
| Journal:           |           |                             |
| □ AIM              | □ BMJ     | □ JAMA                      |
| □ Lancet           | □ NEJM    |                             |

1. Eligible RCT Report?  
   - □ No  
   - □ Exclude → stop here  
     - □ Yes, type of RCT:  
       - □ Two arms  
       - □ Multiple Arms  
       - □ Factorial design

2. Trial described as:  
   - □ Non-inferiority  
   - □ Equivalence  
   - □ Neither

3. Primary outcome clearly specified.  
   - □ Yes, one: ___________________________ (go to q5)  
   - □ No, multiple primary outcomes: ___________________________ (go to q4)  
   - □ None specified (go to q4)

4. If multiple or no primary outcome specified, select one: ______

5. Primary outcome category # (refer to the guide): ___________________________ (e.g. II.3)

6. Effect on primary endpoint reported as:  
   - □ Continuous outcome exclusively  
   - □ Exclude  
     - □ Multinomial outcome exclusively  
     - □ Exclude  
     - □ Binary outcome expressed as rate exclusively  
     - □ Exclude  
     - □ Binary outcome, data not available for 2x2 table  
     - □ Exclude  
     - □ Binary outcome, data available for 2x2 table, go to the next question

7. Is it a composite endpoint?  
   - □ Yes, list components: ___________________________  
   - □ No

8. Is it a patient important outcome?  
   - □ No  
   - □ Exclude  
   - □ Yes, go to the next question

9. Result statistically significant?  
   - □ No  
   - □ Exclude  
   - □ Yes

*Please fill out this box for each study*

- □ Include in FLUKE
- □ Exclude from FLUKE
- □ 3rd reviewer needed (no consensus between 2 reviewers)

**If exclude, reason for exclusion:**  
- □ Not RCT  
- □ Not eligible RCT  
- □ Data for the primary endpoint not available for 2x2 table  
- □ Outcome not patient important  
- □ Result not statistically significant
Data S4 – FLUKE study data abstraction form

FLUKE study: Data Abstraction Form

| Screener initials: | Study ID: | Author, year: |
|--------------------|-----------|---------------|
|                    |           |               |

Journal:  
- [ ] AIM
- [ ] BMJ
- [ ] JAMA
- [ ] Lancet
- [ ] NEJM

1. Eligible RCT Report?  
- [ ] No  
- [ ] Yes, type of RCT:  
  - [ ] Two arms  
  - [ ] Multiple Arms  
  - [ ] Factorial design

2. Trial described as:  
- [ ] Non-inferiority
- [ ] Equivalence
- [ ] Neither

3. Primary outcome clearly specified.  
- [ ] Yes, one: ___________________ (go to q5)  
- [ ] No, multiple primary outcomes: ___________________ (go to q4)  
- [ ] None specified (go to q4)

4. If multiple or no primary outcome specified, select one: ______

5. Primary outcome category # (refer to the guide): ___________________ (e.g. II.3)

6. Effect on primary endpoint reported as:  
- [ ] Continuous outcome exclusively  
- [ ] Exclude
  - [ ] Multinomial outcome exclusively  
  - [ ] Exclude
  - [ ] Binary outcome expressed as rate exclusively  
  - [ ] Exclude
  - [ ] Binary outcome, data not available for 2x2 table  
  - [ ] Exclude
  - [ ] Binary outcome, data available for 2x2 table, go to the next question

7. Is it a composite endpoint?  
- [ ] Yes, list components: ___________________  
- [ ] No

8. Is it a patient important outcome?  
- [ ] No  
- [ ] Exclude
  - [ ] Yes, go to the next question

9. Result statistically significant?  
- [ ] No  
- [ ] Exclude
- [ ] Yes

Please fill out this box for each study

- [ ] Include in FLUKE
- [ ] Exclude from FLUKE
- [ ] 3rd reviewer needed (no consensus between 2 reviewers)

If exclude, reason for exclusion:  
- [ ] Not RCT
- [ ] Not eligible RCT
- [ ] Data for the primary endpoint not available for 2x2 table
- [ ] Outcome not patient important
- [ ] Result not statistically significant
## Background Information

|   |   |
|---|---|
| 10. | Mean/Median Age  
Number of study centers | Age=  
N= |
| 11. | Funding  
*Check all that apply* | ☐ Private only for profit, other  
☐ Private not for profit  
☐ Government  
☐ Not funded  
☐ Not reported |
| 12. | Clinical Area  
*Check only one* | ☐ Medical  
☐ Pharmacological  
☐ Surgical  
☑ Electrophysiology  
☐ Heart failure  
☐ Interventional cardiology  
☐ Open heart surgery  
☐ General cardiology  
☐ Cardiovascular imaging  
☐ Others |
| 13. | Intervention  
*Check only one* | ☐ Pharmacological  
☐ Surgery  
☐ Rehabilitation  
☐ Behavioral intervention  
☐ Complementary and alternative medicine  
☐ Diagnostic test  
☐ Other (specify) |
| 14. | Control  
*Check only one* | ☐ Standard care  
☐ Placebo  
☐ Pharmacological  
☐ Surgery  
☐ Rehabilitation  
☐ Behavioral intervention  
☐ Diagnostic test  
☐ Other (specify) |
### Methodological Quality

| Question                                      | Description                                                                 | Adequate | Inadequate | Not reported |
|-----------------------------------------------|-----------------------------------------------------------------------------|----------|------------|--------------|
| 15. **Concealment of Allocation**             | Check only one                                                              |          |            |              |
|                                               | □ Adequate (involving the use of sequentially numbered, opaque, sealed     |          |            |              |
|                                               |   envelope or coded medication containers or central randomization or      |          |            |              |
|                                               |   quasi-randomized)                                                        |          |            |              |
|                                               | □ Inadequate (Like Open random allocation schedule)                        |          |            |              |
|                                               | □ No method described                                                      |          |            |              |
|                                               | □ Not concealed                                                            |          |            |              |
|                                               | □ Not reported                                                             |          |            |              |
| 16. **Blinding of patients**                  |                                                                             |          |            |              |
|                                               | □ Adequate                                                                  |          |            |              |
|                                               | □ Inadequate                                                                |          |            |              |
|                                               | □ Not reported                                                              |          |            |              |
| 17. **Blinding of health care providers**     |                                                                             |          |            |              |
|                                               | □ Adequate                                                                  |          |            |              |
|                                               | □ Inadequate                                                                |          |            |              |
|                                               | □ Not reported                                                              |          |            |              |
| 18. **Blinding of data collectors**           |                                                                             |          |            |              |
|                                               | □ Adequate                                                                  |          |            |              |
|                                               | □ Inadequate                                                                |          |            |              |
|                                               | □ Not reported                                                              |          |            |              |
| 19. **Blinding of outcome adjudicators**     |                                                                             |          |            |              |
|                                               | □ Adequate                                                                  |          |            |              |
|                                               | □ Inadequate                                                                |          |            |              |
|                                               | □ Not reported                                                              |          |            |              |
| 20. **Blinding of data analysts**             |                                                                             |          |            |              |
|                                               | □ Adequate                                                                  |          |            |              |
|                                               | □ Inadequate                                                                |          |            |              |
|                                               | □ Not reported                                                              |          |            |              |
| 21. **Study stopped early for benefit**       |                                                                             |          |            |              |
|                                               | □ Yes                                                                       |          |            |              |
|                                               | □ No                                                                        |          |            |              |

### ITT Principle

| Question                                      | Description                                                                 | Adequate | Inadequate | Not reported |
|-----------------------------------------------|-----------------------------------------------------------------------------|----------|------------|--------------|
| 22. **Authors used the term ITT**             |                                                                             |          |            |              |
|                                               | □ Yes, ITT                                                                  |          |            |              |
|                                               | □ Yes, Modified ITT                                                         |          |            |              |
|                                               | □ No                                                                        |          |            |              |
| 23. **Post randomization exclusion of         | mistaken by assignment of mistakenly randomized                              |          |            |              |
| mistakenly randomized**                       | □ Yes (Skip Question 24 and 25)                                             |          |            |              |
|                                               | □ No (Go to question 24 and 25)                                             |          |            |              |
|                                               | □ Not reported                                                              |          |            |              |
| 24. **Information about ineligibility was    | available at randomization                                                  |          |            |              |
| available at randomization                    | □ Yes                                                                       |          |            |              |
|                                               | □ No                                                                        |          |            |              |
|                                               | □ Not reported                                                              |          |            |              |
| 25. **Post randomization exclusions were     | blinded to allocation                                                       |          |            |              |
| blinded to allocation                         | □ Yes                                                                       |          |            |              |
|                                               | □ No                                                                        |          |            |              |
|                                               | □ Not reported                                                              |          |            |              |
| 26. **Patients for whom outcome data is       | available were analyzed in the arm to which they were randomized            |          |            |              |
| available were analyzed in the arm to which   | □ Yes                                                                       |          |            |              |
| they were randomized                          | □ No                                                                        |          |            |              |
|                                               | □ Not reported                                                              |          |            |              |
| LTFU statements          |                                                                 |
|-------------------------|-----------------------------------------------------------------|
| 27. LTFU explicitly reported | ☐ Explicit statement: LTFU occurred  
☐ Explicit statement: LTFU did not occur  
☐ No explicit statement about LTFU |
| 28. CONSORT flow diagram | ☐ CONSORT diagram showing LTFU  
☐ CONSORT diagram not showing LTFU  
☐ No CONSORT diagram |
| 29. For studies with no explicit statement about LTFU and no consort diagram | ☐ Meet all 3 prespecified criteria  
☐ Does not meet all 3 prespecified criteria  
☐ N/A |
| 30. LTFU reported separately for the 2 arms | ☐ Yes  
☐ No |
| 31. Authors compared baseline characteristics of LTFU | ☐ Yes  
☐ No |
| 32. Implications of LTFU discussed | ☐ Yes  
☐ No |
| 33. Methods of dealing with LTFU explicitly described | ☐ Yes, methods  
☐ Yes, results  
☐ No |

| Methods of dealing with LTFU |                                                                 |
|-----------------------------|-----------------------------------------------------------------|
| 34. Methods                | Not applicable, no LTFU occurred  
Not applicable, uncertain whether LTFU occurred  
Unclear which method used  
Survival analysis  
Complete case analysis  
Worst case scenario  
Best case scenario  
None of the LTFU had the outcome  
All the LTFU had the outcome  
Different methods for different subgroups of LTFU  
Other (specify) |
| LTFU statistical data | Intervention | control | total | Prespecified assumptions for different groups of LTFU |
|----------------------|--------------|---------|-------|------------------------------------------------------|
| Primary outcome data |              |         |       |                                                      |
| 35. Mistakenly randomized, inappropriately excluded (subtotal 1) |              |         |       |                                                      |
| 36. Did not receive intervention, inappropriately excluded (subtotal 2) |              |         |       |                                                      |
| 37. Withdraw consent (subtotal 3) |              |         |       |                                                      |
| □ unclear whether followed up |              |         |       |                                                      |
| □ not followed up |              |         |       |                                                      |
| □ followed up, not included in the analysis (not LTFU for FLUKE) |              |         |       |                                                      |
| 38. Withdrew consent due to side effect or adverse event |              |         |       |                                                      |
| 39. Withdrew consent due to other specified reason |              |         |       |                                                      |
| 40. Withdrew consent due to unclear reason |              |         |       |                                                      |
| 41. Cross over (subtotal 4) |              |         |       |                                                      |
| □ unclear whether followed up |              |         |       |                                                      |
| □ not followed up |              |         |       |                                                      |
| □ followed up, not included in the analysis (not LTFU for FLUKE) |              |         |       |                                                      |
| □ followed up, analyzed in a group not randomized to (not LTFU for FLUKE) |              |         |       |                                                      |
| 42. Cross over due to side effect or adverse event |              |         |       |                                                      |
| 43. Cross over due to other specified reason |              |         |       |                                                      |
| 44. Cross over due to unclear reason |              |         |       |                                                      |
| 45. Non adherent (subtotal 5) |              |         |       |                                                      |
| □ unclear whether followed up |              |         |       |                                                      |
| □ not followed up |              |         |       |                                                      |
| □ followed up, not included in the analysis (not LTFU for FLUKE) |              |         |       |                                                      |
| □ followed up, analyzed in a group not randomized to (not LTFU for FLUKE) |              |         |       |                                                      |
| 46. Non adherent due to side effect or adverse event |              |         |       |                                                      |
| 47. Non adherent due to other specified reason |              |         |       |                                                      |
| 48. Non adherent due to unclear reason |              |         |       |                                                      |
| 49. Lost contact and no other source of outcome data |              |         |       |                                                      |
|   |   |   |   |
|---|---|---|---|
|50. | Others |   |   |
|51. | LTFU total |   |   |
|   |   |   |   |
|   | LTFU statistical data |   |   |
|   | Primary outcome data | Intervention | control | total |
|52. | Mistakenly randomized, inappropriately excluded (subtotal 1) |   |   |   |
|53. | Did not receive intervention, inappropriately excluded (subtotal 2) |   |   |   |
|54. | Randomized |   |   |   |
|55. | Randomized adjusted (54-53-52) |   |   |   |
|   | Primary outcome data | Intervention events | Control events |   |
|39. | Included in primary analysis |   |   |   |
|40. | Unadjusted effect estimate;95% CI; P value |   |   |   |
Data S5: Further elaboration on the methodology adopted in the systematic review

Analysis method

a. Assessment on the methodological and reporting quality

Bikdeli et al first reported a set of risk factors to consider when evaluating methodological and reporting quality of trials in 2019.14 We consider limiting factors to include the following:

1. Inadequate allocation sequence concealment
2. No blinding on patient
3. Early stoppage
4. Not using intention-to-treat analysis
5. Absence of protocol
6. Without explicit statement on the status of loss to follow up

A univariable random-effects meta-regression analysis was conducted using the log odds of participants lost to follow-up as the dependent variable and general trial characteristics and methodological characteristics as independent variables

1. General trial characteristics
   a. Number of centres
   b. Sample size
   c. Length of follow-up
   d. Type of intervention (Surgery/interventional vs other)
   e. Cardiology Subspecialty (General Medical vs Others)
   f. Type of funding (Commercial Vs Non-profit organisations, governmental or none)

2. Methodological trial characteristics
   a. Concealment of allocation
   b. Blinding of patients
   c. Stopping early for benefit
   d. Use of intention to treat analysis
b. **Extent of loss to follow-up**

The extent of LTFU was estimated by calculating the percentage of LTFU in each trial from each arm (intervention and control). Then, median and interquartile range of the percentages across trials were obtained. A ratio of the total number of participants identified as LTFU to the number of primary outcome events was calculated for each trial (the “lost to follow-up to events ratio”). Median and standard deviation of this ratio was also calculated across the trials.

c. **Potential impact of loss to follow-up**

The potential impact of LTFU is evaluated by proposing assumptions about the outcomes of participants LTFU and the estimated effect of that assumption on the primary outcome (Data S6 for examples). The following common assumptions are first used for calculation:

a. None of the participants lost to follow-up had the event

b. All the participants lost to follow-up had the event

c. None of those lost to follow-up in the treatment group had the event and all those lost to follow-up in the control group did (best case scenario)

d. All participants lost to follow-up in the treatment group had the event and none of those in the control group did (worst case scenario)

Although the above assumptions are widely used in multiple literatures, some experts have countered they are not plausible and have suggested a novel method for estimating effects of LTFU. Akl et al evaluated more plausible assumptions that the incidence of events among participants lost to follow-up is higher by a specific ratio relative to the observed event incidence among participants followed up. They defined $RI_{LTFU/FU}$ as the event incidence among those lost to follow-up relative to the event incidence among those followed up and made plausible assumptions towards the outcome of LTFU participants (see data S6). LTFU refers to “lost to follow-up” and FU refers to “followed up”. A range of plausible $RI_{LTFU/FU}$ values (1,1.5,2,3) was used in both the intervention group and the control group. 3 is the upper limit, which was previously determined by consensus.
Data S6 - Illustrations of the assumptions being considered in FLUKE with examples

Examples based on the following theoretical trial:
- Randomization: 100 to intervention while 100 to control group
- Loss to follow up: 20 in the intervention group while 10 in the control group
- Events: 15 in the intervention group while 20 in the control group

Assumption 1: None of the lost to follow-up participants had the event

|   | intervention | Control |
|---|--------------|---------|
| a | Lost to follow up | 20  | 10 |
| b | Events assumed*  | 0  | 0 |
| c | Events observed in the trial | 15 | 20 |
| d | Total events (b+c) | 15 | 20 |
| e | Randomized | 100 | 100 |
| f | Risk (d/e) | 0.15 | 0.2 |
| g | Relative risk |  | 0.75 |

*None of the lost to follow-up in either group had an event

Assumption 2: All lost to follow-up participants had the event

|   | intervention | Control |
|---|--------------|---------|
| a | Lost to follow up | 20  | 10 |
| b | Events assumed*  | 20  | 10 |
| c | Events observed in the trial | 15 | 20 |
| d | Total events (b+c) | 35 | 30 |
| e | Randomized | 100 | 100 |
| f | Risk (d/e) | 0.35 | 0.3 |
| g | Relative risk | 1.17 |  |

*Each of the lost to follow-up both groups had an event
**Assumption 3: Best case scenario**

|       | intervention | Control |
|-------|--------------|---------|
| a     | Lost to follow up | 20      | 10      |
| b     | Events assumed*  | 0       | 10      |
| c     | Events observed in the trial | 15      | 20      |
| d     | Total events (b+c) | 15      | 30      |
| e     | Randomized     | 100     | 100     |
| f     | Risk (d/e)     | 0.15    | 0.3     |
| g     | Relative risk  | 0.5     |         |

*None of those lost to follow up in the treatment group had the event and all those lost to follow up in the control group did*

**Assumption 4: Worst case scenario**

|       | intervention | Control |
|-------|--------------|---------|
| a     | Lost to follow up | 20      | 10      |
| b     | Events assumed*  | 20      | 0       |
| c     | Events observed in the trial | 15      | 20      |
| d     | Total events (b+c) | 35      | 20      |
| e     | Randomized     | 100     | 100     |
| f     | Risk (d/e)     | 0.35    | 0.2     |
| g     | Relative risk  | 1.75    |         |

*All participants lost to follow up in the treatment group had the event and none of those in the control group did*
Assumptions using relative event incidence (RI_{LTFU/FU})

RI_{LTFU/FU} refers to the event incidence among those lost to follow-up (LTFU) relative to the event incidence among those followed up (FU)

\[
RI_{LTFU/FU} = \frac{\text{Number of events among LTFU}}{\text{number of LTFU}} / \frac{\text{Number of events among FU}}{\text{number of FU}}
\]

- \(RI_{LTFU/FU} = 1\); the event incidence among LTFU and FU is equal
- \(RI_{LTFU/FU} > 1\); the event incidence among LTFU is greater than that of FU

Assumption 1: \(RI_{LTFU/FU} = 1\) in intervention group; and \(RI_{LTFU/FU} = 3\) in control group

|                   | Intervention | Control |
|-------------------|--------------|---------|
| a Lost to follow up | 20           | 10      |
| b Events assumed*  | (20)(1)(15/80)=4 | (10)(3)(20/90)=7 |
| c Events observed in the trial | 15  | 20      |
| d Total events (b+c) | 19  | 27      |
| e Randomized       | 100         | 100     |
| f Risk (d/e)       | 0.19        | 0.27    |
| g Relative risk    |             | 0.70    |

*Number of events assumed = (number lost to follow up) x (RI_{LTFU/FU}) x (Events observed/(number randomized – number lost to follow up))

Assumption 2: \(RI_{LTFU/FU} = 3\) in intervention group; and \(RI_{LTFU/FU} = 1.5\) in control group

|                   | Intervention | Control |
|-------------------|--------------|---------|
| a Lost to follow up | 20           | 10      |
| b Events assumed*  | (20)(3)(15/80)=11 | (10)(1.5)(20/90)=3 |
| c Events observed in the trial | 15  | 20      |
| d Total events (b+c) | 26  | 23      |
| e Randomized       | 100         | 100     |
| f Risk (d/e)       | 0.26        | 0.23    |
| g Relative risk    |             | 1.13    |

*Number of events assumed = (number lost to follow up) x (RI_{LTFU/FU}) x (Events observed/(number randomized – number lost to follow up))
Data S7 – Estimation method accounting for LTFU

Assumptions made using relative event incidence (RI_{LTFU/FU})

RI_{LTFU/FU} refers to the event incidence among those lost to follow-up (LTFU) relative to the event incidence among those followed up (FU) *

Worst RI_{LTFU/FU} assumption = 3 in intervention arm 1 in control arm

Assumption: RI_{LTFU/FU} = 3 in intervention group; and RI_{LTFU/FU} = 1 in control group *

|   | Intervention | Control |
|---|--------------|---------|
| a | Number of Lost to follow up | X | Y |
| b | Event Rate (ER) | (Intervention Event / Number of participants in intervention) | (Control Event/ Number of participants in control) |
| c | Events assumed† | (X)(3)(ER_{inter}) = m | (Y)(1)(ER_{contr}) = n |
| d | Events observed in the trial | Intervention Event | Control Event |
| e | Total events (c+d) | m + Intervention Event | n + Control Event |
| f | Randomized | Number of participants in intervention | Number of participants in control |
| G | Risk (e/f) | R_{inter} | R_{contr} |

*RI_{LTFU/FU} = (Number of events among LTFU / number of LTFU) / (Number of events among FU / number of FU)

- RI_{LTFU/FU} = 1; the event incidence among LTFU and FU is equal
- RI_{LTFU/FU} > 1; the event incidence among LTFU is greater than that of FU

† Number of events assumed = (number lost to follow up) x (RI_{LTFU/FU}) x (Events observed/(number randomized – number lost to follow up))
Data S8 – Among the 23 trials from intervention cardiology, percentage which results would lose significance under different assumptions:

- No events experienced by any lost to follow-up participants =0%
- Events experienced by all the lost to follow-up participants =17%
- Events only experienced by the LTFU in control group while no events experienced by the LTFU in treatment group (best case scenario) =0%
- Events only experienced by the LTFU in treatment group while no events experienced by the LTFU in control group (worst case scenario) =22%

Among the 68 trials from other cardiology field, percentage which results would lose significance under different assumptions:

- No events experienced by any lost to follow-up participants =6%
- Events experienced by all the lost to follow-up participants =9%
- Events only experienced by the LTFU in control group while no events experienced by the LTFU in treatment group (best case scenario) =4%
- Events only experienced by the LTFU in treatment group while no events experienced by the LTFU in control group (worst case scenario) =37%
| Study reference | Country      | Journal     | Mean age |
|-----------------|--------------|-------------|----------|
| S. Verheye (2015) | Belgium      | NEJM        | 67.8     |
| S. S. Anand (2018) | Canada       | The Lancet  | 67.8     |
| M. Dewey (2016)  | Germany      | BMJ         | 60.4     |
| HPS/TIMI55- REVEAL Group (2017) | UK | NEJM        | 67       |
| ASCEND Study Group (2018) | UK | NEJM        | 63.2     |
| H. Calkins (2017) | Germany      | NEJM        | 59.2     |
| C. P. Cannon (2017) | USA | NEJM        | 70.8     |
| Stuart J Connolly (2018) | Canada | The Lancet | 68.3     |
| P J Devereaux (2018) | Canada | The Lancet | 70       |
| J.W. Eikelboom (2018) | Canada | NEJM        | 68.2     |
| R. Estruch (2018) | Spain        | NEJM        | 67       |
| C. M. Gibson (2016) | USA          | NEJM        | 70.1     |
| E. J. Velazquez (2016) | USA | NEJM        | 59.5     |
| J. P. Greenwood (2016) | UK | JAMA        | 56.3     |
| Q. Zhao (2018) | China        | JAMA        | 63.6     |
| B. P Halliday (2018) | UK | The Lancet | 55       |
| A. F. Hernandez (2018) | USA | The Lancet | 64.1     |
| S. C. Johnston (2018) | USA | NEJM        | 65       |
| W.N. Kernan (2016) | USA          | NEJM        | 63.5     |
| JM. Kim (2018) | South Korea  | JAMA        | 60       |
| S. Yusuf (2016) | Canada       | NEJM        | 65.7     |
| N. F. Marrouche (2018) | USA | NEJM        | 64       |
| S. P. Marso (2016) | USA          | NEJM        | 64.3     |
| J. L. Mas (2017)  | France       | NEJM        | 43.7     |
| M. S. Maurer (2018) | USA          | NEJM        | 75       |
| D. E Kandzari (2017) | USA | The Lancet | 64.5     |
| Name                        | Year       | Country     | Journal     | Score |
|-----------------------------|------------|-------------|-------------|-------|
| M.R. Mehra                  | 2018       | USA         | NEJM        | 60    |
| M. R. Mehra                 | 2016       | USA         | NEJM        | 59.6  |
| M.E. Wechsler               | 2017       | USA         | NEJM        | 48.5  |
| A. N Patel                  | 2016       | USA         | The Lancet  | 65    |
| G.D. Perkins                | 2018       | UK          | NEJM        | 69.7  |
| S. R. Steinhubl             | 2018       | USA         | JAMA        | 72.3  |
| P.M. Ridker                 | 2017       | USA         | NEJM        | 61    |
| M. Valgimigli               | 2015       | Netherlands | The Lancet  | 65.8  |
| M. S. Sabatine              | 2017       | USA         | NEJM        | 63    |
| J. L. Sapp                  | 2016       | Canada      | NEJM        | 68.6  |
| J. L. Saver                 | 2017       | USA         | NEJM        | 45.9  |
| G.G. Schwartz               | 2018       | USA         | NEJM        | 58.6  |
| P. C. Smits                 | 2017       | Netherlands | NEJM        | 61.3  |
| B. Zinman                   | 2015       | Canada      | NEJM        | 63.1  |
| L. Søndergaard              | 2017       | Denmark     | NEJM        | 45.2  |
| G.W. Stone                  | 2018       | USA         | NEJM        | 72.3  |
| N. Tegn                     | 2016       | Norway      | The Lancet  | 84.8  |
| H. Thiele                   | 2017       | Germany     | NEJM        | 70    |
| M. Valgimigli               | 2018       | Switzerland | The Lancet  | 65.8  |
| O. Varenne                  | 2017       | France      | The Lancet  | 81.4  |
| S. Yusuf                    | 2016       | Canada      | NEJM        | 65.7  |
| A. Zarbock                  | 2015       | Germany     | JAMA        | 70.4  |
| S.D. Wiviott                | 2018       | USA         | NEJM        | 79.9  |
| M. Abdel-Wahab              | 2014       | Germany     | JAMA        | 80.8  |
| D. H. Adams                 | 2014       | USA         | NEJM        | 83.3  |
| A. Appelboam                | 2015       | UK          | The Lancet  | 54.8  |
| M. P. Bonaca                | 2015       | USA         | NEJM        | 65.3  |
| S. S Brar                   | 2014       | USA         | The Lancet  | 71.5  |
| C. P. Cannon                | 2015       | USA         | NEJM        | 63.6  |
| Author(s)                              | Country     | Journal     | Impact Factor |
|---------------------------------------|-------------|-------------|---------------|
| B. De Bruyne (2014)                   | Belgium     | NEJM        | 63.7          |
| J. D. Douketis (2015)                 | USA         | NEJM        | 71.7          |
| T. Engstrøm (2015)                    | Denmark     | The Lancet  | 63.5          |
| A. M. Gillinov (2015)                 | USA         | NEJM        | 69.6          |
| D. J. Gladstone (2014)                | Canada      | NEJM        | 72.8          |
| THE SPRINT Research Group (2015)      | USA         | NEJM        | 67.9          |
| Y. Han (2015)                         | China       | JAMA        | 57.7          |
| G. Hindricks (2014)                   | Germany     | The Lancet  | 65.5          |
| SJ Hong (2015)                        | Korea       | JAMA        | 64            |
| Y. Huo (2015)                         | China       | JAMA        | 60            |
| M. Imazio (2014)                      | Italy       | The Lancet  | 48.8          |
| M. Imazio (2014)                      | Italy       | JAMA        | 67.5          |
| J. J. McMurray (2014)                 | UK          | NEJM        | 63.8          |
| G. Meyer (2014)                       | Germany     | NEJM        | 66.1          |
| C. A. Morillo (2014)                  | Canada      | JAMA        | 55.3          |
| V. Y. Reddy (2014)                    | USA         | JAMA        | 72            |
| M. Ringh (2015)                       | Sweden      | NEJM        | 72.4          |
| T. Sanna (2014)                       | Italy       | NEJM        | 61.5          |
| A. Shahzad (2014)                     | UK          | The Lancet  | 63.3          |
| P. Urban (2015)                       | Switzerland | NEJM        | 75.7          |
| F. Tomai (2014)                       | Italy       | circulation | 73            |
| M. Valgimigli (2015)                  | Netherlands | JACC        | 71.8          |
| X. D. Zhang (2014)                    | China       | European Heart Journal | 59.2 |
| I. Taguchi (2018)                     | Japan       | circulation | 68.1          |
| Y. D. Tang (2018)                     | China       | circulation | 58.5          |
| J. Zhang (2018)                       | China       | JACC        | 65.6          |
| J. Bermejo (2018)                     | Spain       | European Heart Journal | 71.44 |
| R. S. Bhatia (2017)                   | Canada      | JACC        | 68            |
| D. Bonnet (2017)                      | France      | JACC        | 5.8           |
| Author                  | Year   | Country       | Journal          | Impact Factor |
|------------------------|--------|---------------|------------------|---------------|
| M. Brignole            | 2018   | Italy         | European Heart Journal | 71.5          |
| SL. Chen               | 2017   | China         | JACC             | 64.5          |
| T. Cuisset             | 2016   | France        | European Heart Journal | 60            |
| L. Di Biase            | 2016   | USA           | JACC             | 63.9          |
| L. Di Biase            | 2016   | USA           | circulation      | 61            |
| M. E. Farkouh          | 2018   | Canada        | JACC             | 63.1          |
| P. Garot               | 2016   | France        | JACC             | 75.7          |
| J. P. J. Halcox        | 2017   | UK            | circulation      | 72.6          |
| K. Kaitani             | 2016   | Japan         | European Heart Journal | 63.3          |
| PH Lee                 | 2018   | South Korea   | JACC             | 51.5          |
| G. Sardella            | 2016   | Italy         | JACC             | 72.5          |
| H. Sohara              | 2016   | Japan         | JACC             | 59.5          |
| F. M. Notarangelo      | 2018   | Italy         | JACC             | 70.9          |
| M. Ortiz               | 2017   | Spain         | European Heart Journal | 65.3          |
| J. Yang                | 2014   | China         | JACC             | 5.65          |
| J. Pu                  | 2017   | China         | circulation      | 58            |
| M. Rienstra            | 2017   | Netherlands   | European Heart Journal | 64.5          |
| I. Bernat              | 2014   | Czech Republic | JACC             | 62.1          |
| G. Boriani             | 2014   | Italy         | European Heart Journal | 73.5          |
| D. Carrick             | 2014   | UK            | JACC             | 59.6          |
| A. de Belder           | 2014   | UK            | JACC             | 83.5          |
| L. Di Biase            | 2015   | USA           | JACC             | 66            |
| L. Di Biase            | 2014   | USA           | circulation      | 61.5          |
| A. H. Gershlick        | 2015   | UK            | JACC             | 64.9          |
| Y. Han                 | 2014   | China         | JACC             | 61.4          |
| C. Jennings            | 2014   | UK            | European Heart Journal | 60            |
| J. Layland             | 2015   | UK            | European Heart Journal | 62            |
| M. Leoncini            | 2014   | Italy         | JACC             | 66.2          |
| C. Kaiser              | 2014   | Switzerland   | circulation      | 62.5          |
| Study                  | Country    | Journal                  | Score |
|-----------------------|------------|--------------------------|-------|
| Y. Matsumoto (2014)   | Japan      | JACC                     | 67.5  |
| L. Mont (2014)        | Spain      | European Heart Journal   | 55    |
| G. Montalescot (2014) | France     | European Heart Journal   | 58.2  |
| M. J. Reardon (2015)  | USA        | JACC                     | 83.2  |
Table S2- Sensitivity analysis of the percentage of eligible trials on intervention cardiology vs trials on other subjects in which results would lose significance under different assumptions on the LTFU outcomes on intervention and control group

| Intervention Cardiology † | Others † |
|--------------------------|----------|
| N=23                     | N=68     |
| 3                        | 3        |
| 2                        | 2        |
| 1.5                      | 1.5      |
| 1                        | 1        |

RI LTFU/FU (Control) *

| 1 | 9  | 9  | 0  | 0  | 1  | 1  | 1  | 3  | 6  |
|---|----|----|----|----|----|----|----|----|----|
| 1.5 | 9  | 0  | 0  | 0  | 1.5 | 1  | 3  | 4  | 6  |
| 2  | 9  | 0  | 0  | 0  | 2  | 3  | 5  | 6  | 16 |
| 3  | 0  | 0  | 0  | 0  | 3  | 4  | 12 | 13 | 22 |

N= Number

* RI LTFU/FU is the relative event incidence among those lost to follow-up compared with those followed up

† Paired T test shows that there is significant difference between the subgroup across different assumptions (Mean difference =4.35%, 95%CI 0.295%-8.41%, p=0.0369)
Figure S1 shows the proportion of trials losing significance based on each assumption. It is grouped by the different type of subspecialty. A mean of 3.75% trials form the interventional cardiology subspecialty lost significance while 8.1% trials from other subspecialty lost significance. A paired sample t test was run against the subspecialty yielding a significant difference in proportions (p-value = 0.0369)