Effects of adding ivabradine to usual care in patients with angina pectoris: a systematic review of randomised clinical trials with meta-analysis and Trial Sequential Analysis

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ABSTRACT

Objective To determine the impact of ivabradine on outcomes important to patients with angina pectoris caused by coronary artery disease.

Methods We conducted a systematic review. We included randomised clinical trials comparing ivabradine versus placebo or no intervention for patients with angina pectoris due to coronary artery disease published prior to June 2020. We used Preferred Reporting Items for Systematic Reviews and Meta-Analyses guidelines, Cochrane methodology, Trial Sequential Analysis, Grading of Recommendations Assessment, Development, and Evaluation, and our eight-step procedure. Primary outcomes were all-cause mortality, serious adverse events, and quality of life.

Results We included 47 randomised clinical trials enrolling 35,797 participants. All trials and outcomes were at high risk of bias. Ivabradine compared with control did not have effects when assessing all-cause mortality (risk ratio [RR] 1.04; 95% CI 0.96 to 1.13), quality of life (standardised mean differences −0.05; 95% CI −0.11 to 0.01), cardiovascular mortality (RR 1.07; 95% CI 0.97 to 1.18) and myocardial infarction (RR 1.03; 95% CI 0.91 to 1.16). Ivabradine seemed to increase the risk of serious adverse events after removal of outliers (RR 1.07; 95% CI 1.03 to 1.11) as well as the following adverse events classified as serious: bradycardia, prolonged QT interval, photopsia, atrial fibrillation and hypertension. Ivabradine also increased the risk of non-serious adverse events (RR 1.13; 95% CI 1.11 to 1.16). Ivabradine might have a statistically significant effect when assessing angina frequency (mean difference [MD] 2.06; 95% CI 0.82 to 3.30) and stability (MD 1.48; 95% CI 0.07 to 2.89), but the effect sizes seemed minimal and possibly without any relevance to patients, and we identified several methodological limitations, questioning the validity of these results.

Conclusion Our findings do not support that ivabradine offers significant benefits on patient important outcomes, but rather seems to increase the risk of serious adverse events such as atrial fibrillation and non-serious adverse events. Based on current evidence, guidelines need reassessment and the use of ivabradine for angina pectoris should be reconsidered.

Key questions

What is already known about this subject?

► Ivabradine is recommended in European Society of Cardiology guidelines on chronic coronary syndromes. In the ‘ivabradine for patients with stable coronary artery disease and left-ventricular systolic dysfunction’ (BEAUTIFUL) trial, ivabradine seemed to reduce the incidence of coronary artery disease outcomes in patients with a heart rate of 70 beats/min or higher. In the ‘ivabradine in stable coronary artery disease without clinical heart failure’ (SIGNIFY), ivabradine did not improve clinical outcomes. Previous studies have shown a beneficial effect of ivabradine on angina pectoris symptoms. To our knowledge, no previous systematic review has assessed the effects of ivabradine compared with placebo or no intervention, searching all relevant databases, and considering both risks of systematic errors and random errors.

What does this study add?

► Our findings do not support that ivabradine offers significant benefits on patient important outcomes, but rather seems to increase the risk of serious adverse events such as atrial fibrillation and non-serious adverse events.

How might this impact on clinical practice?

► Based on current evidence, guidelines need reassessment and the use of ivabradine for angina pectoris should be reconsidered.

PROSPERO registration number CRD42018112082.

INTRODUCTION

Cardiovascular diseases accounts for 30% of all deaths worldwide.1 Ischaemic heart disease is associated with an increased risk of mortality and morbidity with an estimated global prevalence over 110 million in 2015.2

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Coronary artery disease is characterised by recurrent episodes of a mismatch between myocardial oxygen supply and demand, resulting in myocardial ischaemia and chest discomfort known as angina pectoris.\(^3\) \(^4\)

Ivabradine is a selective sinus node inhibitor, exerting its effect by decreasing heart rate, thereby decreasing myocardial oxygen demand and increasing myocardial oxygen supply.\(^3\) \(^5\) Theoretically, ivabradine might be an effective intervention for angina pectoris caused by coronary artery disease.\(^6\) \(^7\) To our knowledge, no previous systematic review has assessed the effects of ivabradine compared with usual care (ie, placebo or no intervention) for angina pectoris, searching all relevant databases, and considering both risk of systematic errors and random errors.\(^8\) \(^14\)

**METHODS**

Our methodology is described in detail in our protocol published prior to conducting the literature search.\(^13\) \(^14\)

In short, we carried out this systematic review following the recommendations of the Preferred Reporting Items for Systematic Reviews and Meta-Analyses guidelines.\(^12\)

We included all trials comparing ivabradine versus placebo or no intervention (ie, usual care plus ivabradine vs usual care alone) for participants with angina pectoris. We searched various databases for randomised clinical trials published prior to June 2020, see ‘online supplemental 1’ for a detailed list of databases. Out search strategy is shown in ‘online supplemental 2’. We included randomised clinical trials regardless of trial design, setting, publication status, year, language and reporting of outcomes. Two authors (MM and EEN) independently screened for randomised clinical trials in all non-Chinese databases. The Chinese databases were independently screened by two other authors (LN and SY). Three authors independently extracted data and assessed the risks of bias in the non-Chinese trials (MM all included trials, EEN and NS half each) and two other authors independently extracted data and assessed the risk of bias in the Chinese trials (LN and SY). We attempted to contact trial authors if data were unclear or missing. Disagreements were resolved through discussion or by consulting another author (JCJ).\(^14\)

We assessed three primary outcomes: all-cause mortality, serious adverse events and quality of life. We also assessed three secondary outcomes and eight exploratory outcomes.\(^14\) For all outcomes, we used the trial results reported at maximal follow-up. We chose to assess quality of life using standardised mean differences (SMD), due to the trials reporting on quality of life using different scales. As a ‘rule of thumb’, an effect below 0.4 is a small effect, 0.4 to 0.7 is a moderate effect and above 0.7 is a large effect.\(^15\) \(^16\)

We predefined several subgroup analyses for the primary outcomes, and we conducted sensitivity analyses when assessing both primary and secondary outcomes (see the Results section for sensitivity analyses and see ‘online supplemental material’ for subgroup analyses).\(^14\)

**Assessment of statistical and clinical significance**

We performed all meta-analyses using Review Manager V.5.3.\(^16\) To control for random errors, we used Trial Sequential Analysis and adjusted the threshold for statistical significance as suggested by Jakobsen and colleagues.\(^8\) \(^10\) \(^17\) We used three primary outcomes and therefore considered a p value of 0.025 as the threshold for statistical significance.\(^10\) When analysing secondary and exploratory outcomes, we considered a p value of 0.05 as the threshold for statistical significance.\(^10\) We reported the Trial Sequential Analysis-adjusted confidence intervals and if the cumulative Z-curves crossed any of the Trial Sequential Analysis boundaries (benefit, harm or futility). In order to control the risk of random error when assessing the individual serious and non-serious adverse events, we further adjusted our thresholds for statistical significance according to the large number of comparisons (see ‘Serious adverse events’ and ‘Non-serious adverse events’). Hence, we post-hoc considered a p value of 0.001 as threshold for statistical significance when analysing individual serious and non-serious adverse events.

We used a ‘best-worst case’ and a ‘worst-best case’ analysis to assess the impact of missing data.\(^15\) We used GRADE to assess the certainty of evidence.\(^18\) \(^19\)

**RESULTS**

Our literature search identified 4452 records from databases. We also identified 11 unpublished trials on the trial platform of the company, Servier, that developed ivabradine.\(^20\) After removing duplicates, 3058 records remained. We excluded 2846 records based on title or abstract. We excluded another 165 records based on full text, see ‘online supplemental 3’. We included 42 randomised clinical trials from databases. We included five unpublished, randomised clinical trials from Servier.\(^21\) \(^25\) Therefore, we included a total of 47 randomised clinical trials randomising 35 797 participants.\(^21\) \(^67\) Twenty trials compared ivabradine with placebo,\(^21\) \(^25\) \(^28\) \(^32\) \(^34\) \(^36\) \(^39\) \(^40\) \(^44\) \(^46\) \(^50\) \(^54\) \(^55\) \(^58\) and 27 trials compared ivabradine with ‘no intervention’. Of the 16 trials comparing ivabradine with ‘no intervention’, 14 trials used guideline-based therapy in both trial groups,\(^26\) \(^35\) \(^38\) \(^39\) \(^40\) \(^41\) \(^42\) \(^43\) \(^44\) \(^47\) \(^50\) \(^51\) \(^53\) \(^55\) \(^56\) and 13 trials used various cointerventions other than guideline-based therapy in both trial groups (12 trials used specific beta-blockers,\(^27\) \(^28\) \(^35\) \(^38\) \(^39\) \(^40\) \(^41\) \(^42\) \(^43\) \(^44\) \(^47\) \(^50\) \(^51\) \(^53\) \(^55\) \(^56\) \(^60\) \(^62\) \(^63\) \(^64\) \(^66\) and one used a calcium-channel blocker).\(^33\) For baseline characteristics, see table 1. For all primary and secondary outcomes, we chose to analyse data using fixed-effect meta-analysis due to two trials accounting for more than 97% of weight (see paragraph below).\(^10\) \(^32\) \(^34\)

The two largest trials, the SIGNIFY trial and the BEAUTIFUL trial, contributed with more than 97% of weight.
<table>
<thead>
<tr>
<th>Trial ID</th>
<th>Year</th>
<th>Condition(s)</th>
<th>Max follow-up</th>
<th>Number randomised</th>
<th>Mean age</th>
<th>%-Female</th>
<th>% on beta-blockers</th>
<th>% on calcium channel blockers</th>
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<tr>
<td>Stieg</td>
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<td>Number randomised</td>
<td>Mean age</td>
<td>%-Female</td>
<td>% on beta-blockers</td>
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<td>CAD</td>
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<td>Tagliamonte</td>
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<tr>
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<td>Zhang</td>
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<td>CAD</td>
<td>4 mo</td>
<td>62</td>
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<td>90</td>
<td>61.4</td>
<td>42.2</td>
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AMI, acute myocardial infarction; ASSOCIATE, Efficacy of the I(f) current inhibitor ivabradine in patients with chronic stable angina receiving beta-blocker therapy: a 4-month, randomized, placebo-controlled trial (reference 46); CAD, coronary artery disease; HF, heart failure; mo, months; NR, not reported; RIVENDEL, Heart rate reduction by ivabradine for improvement of endothelial function in patients with coronary artery disease: the RIVENDEL study (reference 41); RIVIERA, Anti-inflammatory effects of ivabradine in patients with acute coronary syndrome: a pilot study (reference 31); wks, weeks.
in all primary and secondary outcome meta-analyses. We identified several methodological limitations regarding these two trials. First, both trials were not prospectively registered before randomisation began. The BEAUTIFUL trial randomised the first participant in January 2005 and was first registered with ClinicalTrials.gov in September 2005 and sent their rationale article for peer review in November 2005. The SIGNIFY trial randomised the first participant in October 2009 and was first registered with ClinicalTrials.gov in May 2015 and first sent their rationale article for peer review in April 2013. Therefore, it was not documented that the methodology, including outcomes and participating centres, was predefined before randomisation began. This is especially problematic when assessing composite outcomes consisting of individual components with very different degrees of severity (ie, in the SIGNIFY trial, the primary composite outcome was death from cardiovascular causes or non-fatal myocardial infarction. In the BEAUTIFUL trial, the primary composite outcome was cardiovascular death, admission to hospital for acute myocardial infarction, or admission to hospital for new onset or worsening heart failure). There is a high risk of selective outcome reporting bias, if the composite outcomes are not clearly predefined before randomisation begins. Furthermore, during the course of the BEAUTIFUL trial, the investigators incorporated a subgroup analysis on participants with a baseline heart rate at or above 70 beats/min (one of the inclusion criteria in the SIGNIFY trial). However, there is no documentation for this subgroup analysis being prespecified prior to initiation of the BEAUTIFUL trial.

Second, in the SIGNIFY trial assessing quality of life and angina pectoris, 13,871 (72.6%) of the 19,102 participants included in the main study were not included in the analysis. It was briefly described in the publication that the reasons were either that some countries did not have a translation of the quality of life scale or it was due to ‘lack of consent’. Third, for serious and non-serious adverse events, there were considerable discrepancies between the data reported in the publication of the SIGNIFY trial as compared with the raw data reported on ClinicalTrials.gov, see ‘online supplemental 11’.

Fourth, both the SIGNIFY trial and the BEAUTIFUL trial, as well as all other included trials, were at high risk of bias. Therefore, there is a risk that our results overestimate beneficial effects and underestimate harmful effects.

We have contacted the trial authors and the company that produced ivabradine, Servier, but we have not received additional information regarding some of these issues. See ‘online supplemental 4’ for a detailed description of the bias risk assessment.

**All-cause mortality**

Fifteen trials randomising 33,427 participants reported on all-cause mortality. Meta-analysis showed no evidence of a difference between ivabradine and control (RR 1.04; 95% CI 0.96 to 1.13; p=0.35; moderate certainty of evidence). Visual inspection of the forest plot and I²-statistics (I²=0%) indicated no heterogeneity. Trial Sequential Analysis showed that we had enough information to reject that ivabradine versus control reduced the risk of all-cause mortality by 15% or more (RR 1.03; 95% CI 0.88 to 1.20; p=0.46; I²=0%; D²=0%). This outcome result was assessed at high risk of bias. Incomplete outcome data alone did not seem to have the potential to influence the results. See ‘Summary of findings-table’ and ‘online supplemental 5’.

**Serious adverse events**

Eighteen trials randomising 33,514 participants reported on serious adverse events. Meta-analysis showed no evidence of a difference between ivabradine and control (RR 1.01; 95% CI 0.98 to 1.04; p=0.56; moderate certainty of evidence). Visual inspection of the forest plot and I²-statistics (I²=55%) indicated substantial heterogeneity which could be resolved by removing the BEAUTIFUL trial and the trial by Tacchetti et al from the analysis (RR 1.06; 95% CI 1.02 to 1.10; p=0.001; I²=0%). Trial Sequential Analysis, after removing outliers, showed that we had enough information to reject that ivabradine decreased the risk of serious adverse events by 15% or more (RR 1.06; 95% CI 1.00 to 1.13; p=0.0014; I²=0%; D²=0%). This outcome result was assessed at high risk of bias. Incomplete outcome data alone did not seem to have the potential to influence the results. See ‘Summary of findings-table’ and ‘online supplemental 6’.

**Individual serious adverse events**

The trials reported on 1,749 different serious adverse events, where the SIGNIFY and the BEAUTIFUL trials reported on the majority of those. To limit problems with multiplicity and type 1 errors caused by the unexpected large number of individual serious adverse events, we post-hoc adjusted the threshold for statistical significance to 0.001 when assessing individual serious adverse events.

Ivabradine increased the risk of the following adverse events classified as serious by the trialists: bradycardia (event classified as ‘bradycardia’ (RR 4.53; 95% CI 2.99 to 6.87; p<0.0001; two trials); event classified as ‘heart rate decreased’ (RR 8.22; 95% CI 3.85 to 17.54; p<0.0001; two trials); event classified as ‘sinus bradycardia’ (RR 6.86; 95% CI 3.11 to 15.15; p<0.0001; one trial)); prolonged QT interval (RR 3.21; 95% CI 1.90 to 5.40; p<0.0001; two trials); photopsia (RR 9.34; 95% CI 2.84 to 30.71; p=0.0002; one trial); atrial fibrillation (RR 1.26; 95% CI 1.10 to 1.44; p=0.0008; three trials); and hypertension (RR 1.42; 95% CI 1.15 to 1.75; p=0.001; one trial).

Ivabradine did not seem to decrease the risk of any individual serious adverse events at the threshold of statistical significance of 0.001.
Quality of life
Four trials randomising 4377 participants reported on quality of life assessed as a continuous outcome. One trial used EuroQoL score (0–100 points), one trial used the Kansas City Cardiomyopathy Questionnaire, one trial used Minnesota Living with Heart Failure Questionnaire, and the SIGNIFY trial reported on quality of life using the Seattle Angina Questionnaire (which does not result in a combined end-score) and a generic visual analogue scale (0–100 points). Meta-analysis showed evidence of a beneficial effect of ivabradine (SMD −0.08; 95% CI −0.14 to −0.02; p=0.009; low certainty of evidence). Visual inspection of the forest plot and I²-statistics (I²=97%) indicated substantial heterogeneity, which could be resolved by removing the trials by Sallam et al and Tatarchenko et al (SMD −0.05; 95% CI −0.11 to 0.01; p=0.09; I²=17%). It was not possible to conduct Trial Sequential Analysis using SMD. This outcome result was assessed as at high risk of bias. We assessed the risk of incomplete outcome data bias to be substantial since 70% of the participants in the SIGNIFY trial were excluded from the analysis of quality of life, see first section of ‘Results’. See ‘Summary of findings-table’ and ‘online supplemental 7’.

Cardiovascular mortality
Eight trials randomising 32 193 participants reported on cardiovascular mortality. Meta-analysis showed no evidence of a difference between ivabradine and control (RR 1.06; 95% CI 0.96 to 1.17; p=0.22; moderate certainty of evidence). Visual inspection of the forest plot and I²-statistics (I²=14%) indicated low heterogeneity. Trial Sequential Analysis showed that we had enough information to reject that ivabradine versus control reduced the risk of cardiovascular mortality by 15% or more (RR 1.066; 95% CI 0.85 to 1.30; p=0.22; I²=14%; D²=53%). This outcome result was assessed at high risk of bias. Incomplete outcome data alone did not seem to have the potential to influence the results. See ‘Summary of findings-table’ and ‘online supplemental 8’.

Myocardial infarction
Five trials randomising 31 810 participants reported on myocardial infarction. Meta-analysis showed no evidence of a difference between ivabradine and control (RR 1.02; 95% CI 0.90 to 1.16; p=0.71; moderate certainty of evidence). Visual inspection of the forest plot and I²-statistics (I²=0%) indicated no heterogeneity. Trial Sequential Analysis showed that we had enough information to reject that ivabradine versus control reduced the risk of myocardial infarction by 15% or more (RR 1.02; 95% CI 0.85 to 1.23; p=0.71; I²=0%; D²=0%). This outcome result was assessed at high risk of bias. Incomplete outcome data alone did not seem to have the potential to influence the results. See ‘Summary of findings-table’ and ‘online supplemental 9’.

Non-serious adverse events
Twenty-four trials randomising 34 181 participants reported on non-serious adverse events. Meta-analysis showed evidence of a harmful effect of ivabradine (RR 1.13; 95% CI 1.11 to 1.16; p<0.0001; moderate certainty of evidence). Visual inspection of the forest plot and I²-statistics (I²=8%) indicated low heterogeneity. Trial Sequential Analysis showed that we had enough information to detect that ivabradine versus control increased the risk of non-serious adverse events by 13% or more (RR 8.05; 95% CI 6.76 to 9.59; p<0.0001; two trials); event classified as ‘bradycardia’ (RR 9.61; 5.65 to 16.33; p<0.0001; one trial); QT interval prolonged (RR 2.65; 95% CI 1.85 to 3.81; p<0.0001; one trial); hypertension (RR 1.19; 95% CI 1.09 to 1.30; p=0.0001; four trials); and dizziness (RR 1.32; 95% CI 1.13 to 1.54; p=0.0003; four trials).

Individual non-serious adverse events
The trials reported 54 different non-serious adverse events; the SIGNIFY and the BEAUTIFUL trials reported on the majority of those. To limit problems with multiplicity and type 1 errors caused by the unexpected large number of individual serious adverse events, we post-hoc adjusted the threshold for statistical significance to 0.001 when assessing individual non-serious adverse events.19

Ivabradine seemed to increase the risk of bradycardia (event classified as ‘bradycardia’ (RR 4.54; 95% CI 3.78 to 5.46; p<0.0001; nine trials); event classified as ‘heart rate decreased’ (RR 8.05; 95% CI 6.76 to 9.59; p<0.0001; two trials); event classified as ‘sinus bradycardia’ (RR 9.61; 5.65 to 16.33; p<0.0001; one trial)); phosphenes (RR 6.58; 95% CI 5.34 to 8.10; p<0.0001; seven trials); vision blurred (RR 3.39; 95% CI 2.32 to 4.93; p<0.0001; five trials); QT interval prolonged (RR 2.65; 95% CI 1.85 to 3.81; p<0.0001; one trial); and dizziness (RR 1.32; 95% CI 1.13 to 1.54; p=0.0003; four trials).

Ivabradine seemed to decrease the risk of sinus tachycardia (RR 0.25; 95% CI 0.18 to 0.35; p=0.0001; one trial).

Exploratory outcomes
Ivabradine seemed to increase the score of angina frequency (0–100 points, higher score represents a positive outcome; MD 2.06; 95% CI 0.82 to 3.30; p=0.001; I²=0%; 3 trials; 4297 participants) and angina stability, both on the Seattle Angina Questionnaire score (0–100 points, higher score represents a positive outcome; MD 1.48; 95% CI 0.07 to 2.89; p=0.04; I²=0%; 2 trials; 4217 participants). However, the effect sizes were minimal (SMD 0.1 for angina frequency and SMD 0.06 for angina stability) and both were more than five times below our predefined minimal important difference (SMD 0.5). Furthermore, the SIGNIFY trial accounted for more than 97% of the total weight in both analyses and used questionable methodology (see the first section of the Results section). We assessed the risk of incomplete outcome data bias to be substantial since 70% of the participants in the SIGNIFY trial were excluded from the analysis of
angina, see the first section of the Results section and ‘online supplemental 11’.

Four trials assessed exercise tolerance tests. In the ASSOCIATE trial, ivabradine versus control did not reach the minimal important difference in any of the outcome measures in exercise tolerance testing (time to angina onset, time to limiting angina, time to 1 mm ST depression and total exercise duration). In the trial by Borer et al., ivabradine treatment reached the minimal important difference for time to angina onset and time to 1 mm ST depression in the 10 ng ivabradine twice daily group, which is a higher dose than recommended. The participants had discontinued any other anti-ischaemic drug two to 7 days prior to randomisation and the primary outcome was measured after 14 days of treatment. In two of the unpublished trials, there seemed to be no difference between ivabradine and control in regard to exercise tolerance testing.

The results of the remaining exploratory outcomes are reported in the online supplemental material, see ‘online supplemental 12’ and the Discussion section.

Subgroup analyses
We post-hoc decided to conduct a subgroup analysis of trials randomising participants with coronary artery disease alone compared to trials randomising participants with both coronary artery disease and heart failure with reduced ejection fraction. We identified 11 trials including participants with both coronary artery disease and heart failure with an ejection fraction of 40% or less and one trial including participants with an ejection fraction of 45% or more. We judged trials as being ‘coronary artery disease only’ trials, if heart failure was not an inclusion criterion, if heart failure was an exclusion criterion, or if there was no mention of heart failure. For serious adverse events, test for subgroup differences showed evidence of a difference (p<0.00001). When analysed separately, we found evidence of a harmful effect of ivabradine in trials randomising participants with coronary artery disease only (RR 1.06; 95% CI 1.02 to 1.10; p=0.001; I²=0%; four trials) and evidence of a beneficial effect of ivabradine in trials administering ivabradine below median daily dose (RR 0.92; 95% CI 0.87 to 0.97; p=0.002; I²=26%; four trials).

For the remaining subgroup analyses, test for subgroup differences showed no evidence of a difference between ivabradine and control. The results of all subgroup analyses can be found in ‘online supplemental 5–7’.

DISCUSSION
We included a total of 47 randomised clinical trials enrolling 35 797 participants. All trials and outcomes were at high risk of bias.

Meta-analyses and Trial Sequential Analysis showed that there was no evidence of a difference between ivabradine and placebo or no intervention when assessing all-cause mortality, quality of life, cardiovascular mortality and myocardial infarction.

Meta-analysis showed that ivabradine potentially increased the risk of serious adverse events after removal of outliers from the analyses as well as the following adverse events classified as serious: bradycardia, prolonged QT interval, photopsia, atrial fibrillation and hypertension. Meta-analysis and Trial Sequential Analysis showed that ivabradine seemed to increase the risk of non-serious adverse events.

Ivabradine seemed to increase the score of angina frequency and angina stability on the Seattle Angina Questionnaire. However, we identified several methodological limitations regarding these outcomes (see the Results section). Furthermore, the observed effect sizes seemed minimal as they were more than five times lower than our predefined minimal important difference.

See ‘Summary of findings-table’ in ‘online supplemental 13’.

In a post-hoc subgroup analyses, ivabradine seemed to increase the risk of serious adverse events in trials randomising participants with only coronary artery disease and to decrease the risk of serious adverse events in trials randomising participants with both coronary artery disease and heart failure. Post-hoc analyses need to be interpreted with caution.

Our systematic review has several strengths. Our methodology was predefined and was described in detail in our published protocol. We identified 47 trials, whereas the largest of the previous (non-systematic) reviews only included eight trials. To control the risk of random error, we used Trial Sequential Analysis and adjusted our thresholds for statistical significance. To control the risk of systematic error, we assessed the risk of bias of all included trials. To assess if the thresholds for statistical and clinical significance were crossed, we used our eight-step procedure. We included all randomised clinical trials regardless of publication type, status, language and outcomes. We attempted to...
contact trialists if there were incomplete outcome data or additional information was required.

Our review also has several limitations. All trials were at high risk of bias, including a substantial risk of for-profit bias.40 Fourteen of the trials were sponsored by the company that developed ivabradine, including the two largest trials, the SIGNIFY and the BEAUTIFUL trials, that randomised 30,019 participants (91%).21–25 Sponsorship of drug trials by manufacturing companies leads to more favourable efficacy results than trials sponsored by other sources.80 The BEAUTIFUL and the SIGNIFY trials both used composite outcomes that did not seem to be predefined prior to randomising participants, see the Results section.32 54 70–73 The results of 11 trials were reported only as abstracts, which made the assessment of the methodology and results problematic.7 There is a risk that our present results overestimate the beneficial effects and underestimate the harmful effects of ivabradine.80–82 Last, to limit problems with multiplicity and type I errors caused by the unexpected large number of individual serious adverse events, we post-hoc adjusted the threshold for statistical significance to 0.001 when assessing individual serious and non-serious adverse events. This threshold was not predefined, and these results should therefore be interpreted with caution. Nevertheless, several of the harmful outcomes may indeed be increased by ivabradine.80–82

Ivabradine was recommended as an effective second-line treatment for angina relief in the 2013 and the 2019 European Society of Cardiology guidelines on the management of chronic coronary artery syndromes.3 82 However, we did not identify any valid evidence supporting that ivabradine should reduce angina symptoms in a clinically significant way. On the contrary, our results show that ivabradine does not seem to have beneficial effects on all-cause mortality, serious adverse events, quality of life, cardiovascular mortality and myocardial infarction. Moreover, ivabradine increases the risk of both serious and non-serious adverse events. The lack of benefit has previously been discussed as being partly due to the decrease in heart rate caused by ivabradine resulting in an increase in systolic blood pressure and left ventricular overload.93

CONCLUSION
Our findings do not support that ivabradine offers significant benefits on patient important outcomes, but rather seems to increase the risk of serious adverse events such as atrial fibrillation and non-serious adverse events. Based on current evidence, guidelines need reassessment and the use of ivabradine for angina pectoris should be reconsidered.

Differences between protocol and review
We conducted our literature search in parallel with another review on the effects of adding ivabradine to usual care in participants with heart failure. We originally planned to report and analyse the results including participants with angina and participants with heart failure in one review, but due to clinical and statistical heterogeneity and reviewer recommendations, we decided to report the results in two separate reviews.14

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