Morphine in acute coronary syndrome: systematic review and meta-analysis

Gonçalo Silva Duarte, Afonso Nunes-Ferreira, Filipe Brogueira Rodrigues, Fausto J Pinto, Joaquim J Ferreira, Joao Costa, Daniel Caldeira

ABSTRACT

Objective Morphine is frequently used in acute coronary syndrome (ACS) due to its analgesic effect, it being recommended in the main cardiology guidelines in Europe and the USA. However, controversy exists regarding its routine use due to potential safety concerns. We conducted a systematic review of randomised-controlled trials (RCTs) and observational studies to synthesise the available evidence.

Design Systematic review and meta-analysis.

Data sources CENTRAL, MEDLINE, EMBASE and trial registries.

Eligibility criteria for selecting studies We included RCTs and observational studies evaluating the impact of morphine in cardiovascular outcomes or platelet reactivity measures.

Data extraction and synthesis Data were screened, extracted and appraised by two independent reviewers. The data were pooled results using a random-effects model. Outcomes included in-hospital mortality, major adverse cardiovascular events (MACE), platelet reactivity (using VerifyNow) and bleeding, reported as relative risk (RR) with 95% CI. We assessed the confidence in the evidence using the Grading of Recommendations Assessment, Development and Evaluation (GRADE) framework. We followed the Meta-Analysis Of Observational Studies in Epidemiology and Preferred Reporting Items for Systematic Reviews and Meta-Analyses guidelines.

Results Five RCTs and 12 observational studies were included, enrolling 69,993 participants. Pooled results showed an increased risk of in-hospital mortality (RR 1.45 [95% CI 1.10 to 1.91], low GRADE confidence), MACE (RR 1.21, 95% CI 1.02 to 1.45) and an increased platelet reactivity at 1 and 2 hours (99.37 platelet reactivity units [PRU], 95% CI 36.04 to 82.71; 68.28 PRU, 95% CI 37.01 to 99.55, high GRADE confidence) associated with morphine. We found no significant difference in the risk of bleeding. We found no differences in subgroup analyses based on study design and ACS subtype.

Conclusions Morphine was associated with an increased risk of in-hospital mortality and MACE but the high risk of bias leads to low result confidence. There is high confidence that morphine decreases the antiplatelet effect of P2Y12 inhibitors.

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INTRODUCTION

Worldwide, cardiovascular events are the leading cause of death.1 The burden of disease will likely remain high2 3 as the incidence of cardiovascular events is expected to continue increasing.4

Antiplatelet agents (aspirin and P2Y12 inhibitors), anticoagulants and coronary revascularisation are the mainstay in the early treatment of acute coronary syndrome (ACS), and their use likely improves prognosis.5 7 In both Europe and the USA, morphine, a potent analgesic that is a competitive agonist of μ-receptors in the central nervous system and smooth muscle, is recommended for pain control in the ACS setting.5 However, the activation of opioid receptors in the myenteric plexus decreases gut motility and secretion, inhibiting the activation of P2Y12 inhibitors by decreasing their absorption and bioavailability.6 There is conflict regarding the possibility that morphine interferes with P2Y12 inhibitors in the achievement of an adequate antithrombotic milieu,7 9 10 which may decrease the efficaciousness of antiplatelet drugs, if these are given concurrently with morphine. We conducted a systematic review of randomised-controlled trials (RCTs) and observational studies to evaluate the safety of morphine use in ACS, hypothesising that we would find a clinically meaningful result.
METHODS
This systematic review followed the reporting principles of Meta-analysis Of Observational Studies in Epidemiology and Preferred Reporting Items for Systematic Reviews and Meta-Analyses. Patients and public were not involved in this review.

Eligibility criteria
We considered longitudinal studies (ie, RCTs and observational studies) evaluating the impact of morphine in cardiovascular outcomes or platelet reactivity measures. The target population was patients with ACS, which can be either ST-elevated myocardial infarction (STEMI) or non-ST elevated acute coronary syndrome (NSTE-ACS). Studies had to evaluate morphine (irrespective of the administration route or dose) against placebo, control (no intervention arm) or any other analgesic non-opioid drug.

Primary outcomes were in-hospital mortality and major adverse cardiovascular events (MACE), as defined by the PLATO trial (cardiovascular mortality, non-fatal MI or non-fatal stroke). Secondary outcomes comprised additional safety outcomes, as defined within the included studies (as reported in the original studies, including morphine-related adverse events such as bleeding, nausea/ emesis, bradycardia, hypotension and respiratory insufficiency) and platelet reactivity (a pharmacodynamic outcome, sought through the VerifyNow method, which is the most widely used assay to evaluate platelet reactivity and shows a stronger correlation with MACE in ACS than other methods, namely multiple electrode aggregometry (MEA)/Multiplate). VerifyNow is a blood test that measures platelet reactivity by the rate and extent of light changes in whole blood as platelets aggregate, and therefore measures platelet response to major antiplatelet agents.

Information sources and search method
Potentially eligible studies were identified through an electronic search of CENTRAL (Cochrane Library), MEDLINE, EMBASE, and clinicaltrials.gov from inception to November 2018 (online supplementary material). No language restrictions were applied. We cross-checked reference lists of reports for potential additional studies.

Study selection and data collection process
Two reviewers (GSD and either FBR or ANF) independently screened the titles and abstracts yielded by the search and assessed the full texts of the selected studies to determine the appropriateness for inclusion. Disagreements were resolved through consensus or by a third reviewer (DC) serving as final arbitrator. The reasons for exclusion were recorded at the full-text screening stage.

Two reviewers (ANF and GSD) extracted study data following a pre-established data collection form. Data from studies’ plots were retrieved through Plot Digitizer V.2.6.8. When studies presented different estimates of the outcome of interest, we extracted the most precise or adjusted measures.

Risk of bias was independently evaluated by two authors (GSD and ANF) using different tools according to study design. For RCTs, we used the Cochrane risk of bias tool, where domains were qualitatively classified as at high, unclear or low risk of bias. The overall risk of bias for each RCT was divided as high or low risk, with high risk being those RCTs in which at least one domain was assessed at a high risk of bias, or more than three domains were had a rating of unclear. For observational studies we used the ROBINS-I tool, assessing the following domains: confounding, selection of participants, classification of intervention, deviations from intervention, missing data, measurement of outcome and selection of reported results. These domains were qualitatively classified as at critical, serious, moderate or low risk of bias. The overall risk of bias for each observational study was divided as critical or non-critical, following ROBINS-I criteria. Risk of bias graphs was derived from these tools.

Statistical analysis
We used OpenMetaAnalyst and Review Manager for statistical analysis and to derive forest plots. We used a random-effects model to pool data owing to the anticipated heterogeneity in the included trials, in particular differences in study design. We reported pooled dichotomous data using risk ratios (RRs) and continuous data the mean difference (MD), reporting 95% CIs and corresponding p values for both. Heterogeneity was assessed using I². When present effect estimates as RR because relative estimates are more similar across studies with different designs, populations and lengths of follow-up than absolute effects. When raw data or RR was unavailable, we used the HR or OR provided the estimate was small. Preplanned subgroup analyses considering study design (RCTs and observational studies) and ACS type were conducted. A sensitivity analysis was also performed, in which RCTs at high risk of bias and observational studies at critical risk of bias were excluded from the analysis. Reporting bias was performed through funnel plot examination and statistical methods providing that a sufficient number of studies were included.

When observational studies were assessed as having a critical risk of bias, we adjusted the within-study variance– covariance matrix using a precision correction of 0.1 that will provide more conservative pooled estimates. This conservative weight factor was based on expert-based clinical grounds.

We used the Grading of Recommendations, Assessment and Evaluation (GRADE) framework to report the overall quality of evidence. The certainty in the evidence for each outcome was graded as high, moderate, low or very low.

Patient and public involvement
Patients and the public were not involved in this review.

**RESULTS**

**Included studies**

The search returned 1419 records, resulting in 1035 records after removing all duplicates. After title and abstract screening, 53 articles were assessed for full-text screening, with 17 being included for qualitative and quantitative syntheses, 5 being RCTs and 12 being observational studies. We did not retrieve any unpublished study (figure 1).

The characteristics of the included studies can be seen in table 1 and table 2. Study publication dates ranged from 1969 to 2018, with sample sizes between 12 and 57039 participants. The largest study, Meine et al., a retrospective cohort study, accounted for 81% of the participants in this review.

Morphine administration was variable across the included studies, with six of the observational studies not reporting information regarding dose and number or mode of administration. Among RCTs, morphine administration was intravenous or intramuscular at a dose between 2 and 10 mg, either in single or multiple administrations. The forms of antiplatelet therapy used across studies were varied, and firm conclusions cannot be made.

**Risk of bias**

We judged two of the five RCTs to be at a high overall risk of bias, one due to having unclear risk of bias in all but one domain, and the other due to a high risk of performance and attrition bias. All observational studies were at risk of bias due to confounding, and all but one were at moderate risk of selection of study results. Grendahl and Hansteen38 was additionally at moderate risk of bias due to measurement of outcome. Overall, two observational studies were at critical risk of bias, seven were at serious risk of bias and three were at a moderate risk of bias (online supplementary material).

**In-hospital mortality**

Four RCTs (n=2237) and seven observational studies (n=63112) contributed with data for this outcome. Adjusted pooled results showed an increased risk of in-hospital mortality in the morphine group (RR 1.45; 95% CI 1.10 to 1.91; I²=0%; figure 2). Subgroup analysis
Table 1  Characteristics of the randomised controlled trials

<table>
<thead>
<tr>
<th>Study</th>
<th>Location</th>
<th>Mean follow-up</th>
<th>Patients</th>
<th>Antiplatelet medication used</th>
<th>Morphine characteristics</th>
<th>Comparator</th>
<th>N (total)</th>
<th>Mean age</th>
<th>Primary outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bressan et al</td>
<td>Single centre, Italy</td>
<td>24 hours</td>
<td>Patients with AMI, chest pain and symptoms&lt;6 hour</td>
<td>None</td>
<td>10mg IM single dose</td>
<td>Indoprofen</td>
<td>40</td>
<td>54</td>
<td>Assessment of analgesic effect of indoprofen in AMI patients</td>
</tr>
<tr>
<td>Everts et al</td>
<td>Single centre, Sweden</td>
<td>6 months</td>
<td>Patients admitted to the coronary care unit because of symptoms of suspected AMI</td>
<td>None</td>
<td>2–7.5 mg IV, single to multiple doses</td>
<td>Metoprolol</td>
<td>265</td>
<td>66.6</td>
<td>Assessment of analgesic effect of metoprolol in suspected or definitive AMI patients</td>
</tr>
<tr>
<td>Kubica et al</td>
<td>Single centre, Poland</td>
<td>Hospital stay</td>
<td>Patient with the diagnosis of STEMI or NSTEMI</td>
<td>Aspirin and ticagrelor</td>
<td>5mg IV single dose</td>
<td>Placebo</td>
<td>70</td>
<td>61.6</td>
<td>Assess the influence of morphine on pharmacokinetics and pharmacodynamics of ticagrelor and its active metabolite</td>
</tr>
<tr>
<td>Lapostolle et al</td>
<td>Multicentre</td>
<td>30 days</td>
<td>STEMI</td>
<td>Aspirin and ticagrelor</td>
<td>NR</td>
<td>Placebo</td>
<td>1862</td>
<td>60.8</td>
<td>TIMI Flow Grade 3 of culprit vessel at initial angiography and ST-segment elevation resolution Pre-PCI≥70% (coprimary outcomes)</td>
</tr>
<tr>
<td>Thomas et al</td>
<td>Single centre, UK</td>
<td>24 hours</td>
<td>STEMI</td>
<td>Aspirin and prasugrel</td>
<td>5mg IV single dose</td>
<td>Placebo</td>
<td>12</td>
<td>64</td>
<td>VerifyNow platelet reactivity</td>
</tr>
</tbody>
</table>

AMI, acute myocardial infarction; IM, intramuscular; IV, intravenous; NR, not reported; NSTEMI, non-ST elevation myocardial infarction; PCI, percutaneous coronary intervention; STEMI, ST elevation myocardial infarction; TIMI, thrombolysis in myocardial infarction.
<table>
<thead>
<tr>
<th>Study</th>
<th>Location, study design</th>
<th>Follow-up (years)</th>
<th>Patients</th>
<th>Antiplatelet medication used</th>
<th>Morphine characteristics</th>
<th>Comparator</th>
<th>N</th>
<th>Mean age (SD)</th>
<th>Outcome measures</th>
<th>Ascertainment</th>
<th>Outcome adjustments for confounders</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bellandi et al.</td>
<td>Italy, Greece, prospective</td>
<td>2</td>
<td>STEMI patients undergoing PCI and receiving either prasugrel or ticagrelor</td>
<td>NR</td>
<td>6–3 mg, no additional information</td>
<td>No intervention</td>
<td>182</td>
<td>64 (13)</td>
<td>Myocardial reperfusion by early ST-segment resolution</td>
<td>According to the physician’s decision</td>
<td>Operator blinded to morphine use</td>
</tr>
<tr>
<td>Bonin et al.</td>
<td>France, retrospective</td>
<td>1</td>
<td>STEMI</td>
<td>NR</td>
<td>NR</td>
<td>No intervention</td>
<td>969</td>
<td>60 (13)</td>
<td>MACE</td>
<td>According to the physician’s decision</td>
<td>NR</td>
</tr>
<tr>
<td>Danchin et al.</td>
<td>France, retrospective</td>
<td>1</td>
<td>STEMI</td>
<td>DAPT</td>
<td>NR</td>
<td>No intervention</td>
<td>3548</td>
<td>63 (12)</td>
<td>All-cause mortality</td>
<td>According to the physician’s decision</td>
<td>NR</td>
</tr>
<tr>
<td>Farag et al.</td>
<td>UK, prospective single centre</td>
<td>30 days</td>
<td>STEMI</td>
<td>DAPT</td>
<td>5–10 mg IV</td>
<td>No intervention</td>
<td>300</td>
<td>64 (13)</td>
<td>MACE and major bleeding</td>
<td>According to the physician’s decision</td>
<td>NR</td>
</tr>
<tr>
<td>Franchi et al.</td>
<td>USA, posthoc analysis of RCT</td>
<td>1</td>
<td>STEMI patients undergoing PCI</td>
<td>Aspirin and ticagrelor</td>
<td>NR</td>
<td>No intervention</td>
<td>46</td>
<td>59</td>
<td>Asses the pharmacokinetic and pharmacodynamics of escalating doses of ticagrelor</td>
<td>According to the physician’s decision</td>
<td>VerifyNow and VASP assays</td>
</tr>
<tr>
<td>Grendahland and Hansteen</td>
<td>Norway, prospective</td>
<td>NR</td>
<td>Uncomplicated AMI&lt;48hour of symptoms</td>
<td>NR</td>
<td>150mg IV/single dose</td>
<td>Placebo</td>
<td>20</td>
<td>NR</td>
<td>Assess the circulatory effects of morphine</td>
<td>NR</td>
<td>NR</td>
</tr>
<tr>
<td>Johnson et al.</td>
<td>UK, posthoc analysis</td>
<td>1.5</td>
<td>STEMI</td>
<td>Aspirin and prasugrel</td>
<td>NR</td>
<td>No intervention</td>
<td>106</td>
<td>61.1 (11.7)</td>
<td>Platelet reactivity</td>
<td>According to the physician’s decision</td>
<td>Multiplate assay</td>
</tr>
<tr>
<td>McCarthy et al.</td>
<td>USA, retrospective</td>
<td>Hospital stay</td>
<td>STEMI and NSTEMI-ACS</td>
<td>Insufficient detail</td>
<td>NR</td>
<td>No intervention</td>
<td>3027</td>
<td>62 (12)</td>
<td>Mortality</td>
<td>According to the physician’s decision</td>
<td>NR</td>
</tr>
<tr>
<td>Meine et al.</td>
<td>USA, retrospective</td>
<td>2.5</td>
<td>NSTEMI</td>
<td>Insufficient detail</td>
<td>IV, no additional information</td>
<td>No intervention</td>
<td>57 039</td>
<td>68</td>
<td>Assess in-hospital death, recurrent myocardial infarction, congestive heart failure, cardiogenic shock</td>
<td>According to the physician’s decision</td>
<td>CRUSADE database</td>
</tr>
<tr>
<td>Puymirat et al.</td>
<td>France, Retrospective</td>
<td>2months</td>
<td>STEMI patients with symptoms&lt;48hour</td>
<td>50% Aspirin</td>
<td>71% Clopidogrel</td>
<td>8% Prasugrel</td>
<td>NR</td>
<td>No intervention</td>
<td>2438</td>
<td>63</td>
<td>Assess practices for myocardial infarction management in ‘real life’ and with medium and long-term outcomes</td>
</tr>
<tr>
<td>Siller-Matula et al.</td>
<td>Austria, prospective</td>
<td>2</td>
<td>STEMI patients treated with in-hospital loading dose of prasugrel</td>
<td>Aspirin and prasugrel</td>
<td>5–15mg IV single dose</td>
<td>No intervention</td>
<td>32</td>
<td>60 (11)</td>
<td>Assess if abciximab is a bridging therapy to achieve adequate levels of platelet inhibition</td>
<td>According to the physician’s decision</td>
<td>Multiplate assay</td>
</tr>
<tr>
<td>Silvain et al.</td>
<td>International, posthoc analysis of RCT</td>
<td>14hours</td>
<td>STEMI</td>
<td>Aspirin and ticagrelor</td>
<td>NR</td>
<td>No intervention</td>
<td>37</td>
<td>56.2 (10.2)</td>
<td>Assess coronary reperfusion prior to percutaneous coronary intervention with prehospital or in-hospital ticagrelor 180mg loading dose</td>
<td>According to the physician’s decision</td>
<td>VerifyNow assay</td>
</tr>
</tbody>
</table>

AMI, acute myocardial infarction; DAPT, dual antiplatelet therapy; IV, intravenous; MACE, major adverse cardiovascular events; NR, not reported; NSTEMI, non-ST elevation myocardial infarction; PCI, primary percutaneous coronary intervention; RCT, randomised controlled trial; STEMI, ST elevation myocardial infarction; VASP, vasodilator-stimulated phosphoprotein.
based on study design (p=0.67 for interaction; figure 2) and ACS subtype (STEMI RR 1.05; 95% CI 0.57 to 1.94; I²=0%; NSTE-ACS RR 1.57; 95% CI 1.15 to 2.14; I²=0% and p=0.25 for interaction) were both non-significant. Sensitivity analysis by excluding studies at critical risk of bias showed no differences between morphine and control (RR 1.41; 95% CI 0.87 to 2.27; I²=0%; n=5872 participants). The GRADE confidence in this estimate is low.

**MACE**

Three RCTs (n=375) and seven observational studies (n=61,054) contributed with data for this outcome. Adjusted pooled results showed an increased risk of MACE in the morphine group (RR 1.21, 95% CI 1.02 to 1.45; I²=0%; figure 3). Subgroup analysis based on study design (p=0.44 for interaction; figure 3) and ACS subtype (STEMI RR 1.20; 95% CI 0.71 to 2.03; I²=0%; NSTE-ACS RR 1.21; 95% CI 1.01 to 1.46; I²=0% and p=0.98 for interaction) were both non-significant. Sensitivity analysis by excluding studies at critical risk of bias showed no differences between morphine and control (RR 1.40; 95% CI 0.85 to 2.30; I²=0%; n=1952). The GRADE confidence in this estimate is low.

**Platelet reactivity**

We present data from 1 and 2 hours after morphine administration, as these are likely to be the most clinically meaningful timepoints. Two RCT (n=82) and two observational studies (n=228) contributed with data for this outcome. One hour after administration, morphine was associated with increased platelet reactivity, with an MD of 59.37 platelet reactivity units (PRU) (95% CI 36.04 to 82.71; I²=23%; figure 4). Two hours after administration, morphine remained associated with increased platelet reactivity (MD 68.28 PRU, 95% CI 37.01 to 99.55; I²=28%; figure 5). Subgroup analysis based on study design and ACS subtype were both non-significant at both timepoints (p=0.25 for interaction for both timepoints; p=0.24 for interaction for both timepoints, respectively). The GRADE confidence is high for this outcome at both timepoints.

We additionally pooled results using the three trials that reported results using the MEA method. These results were consistent with those using the VerifyNow
method at both 1 hour (MD 27.80, 95% CI 16.03 to 39.57, I²=24%) and 2 hours after morphine administration (MD 19.99, 95% CI 1.52 to 38.46, I²=82%).

Additional outcomes

We found no differences associated with morphine use, namely regarding the risk of cardiogenic shock (RR 1.48; 95% CI 1.00 to 2.18; I²=0%), heart failure (RR 1.17; 95% CI 0.91 to 1.51; I²=33%), hypotension (RR 0.93; 95% CI 0.49 to 1.74; I²=5%), nausea/emesis (RR 1.84; 95% CI 0.80 to 4.23; I²=44%), respiratory insufficiency (RR 0.77; 95% CI 0.31 to 1.91; I²=0%) or stent thrombosis (RR 1.13; 95% CI 0.67 to 1.92; I²=0%) (online supplementary material).

DISCUSSION

Our main findings were as follows: (1) morphine was associated with an increased risk of in-hospital mortality and MACE; however, high risk of bias led to low confidence in the results; (2) morphine decreased the antiplatelet effect of P2Y12 inhibitors in the first hours of ACS, and

![Figure 3](https://example.com/figure3.png)

**Figure 3** Forest plot of MACE (major adverse cardiovascular events) according to morphine use, subgroups according to study design. IV, inverse variance; RCTs, randomised controlled trials.

![Figure 4](https://example.com/figure4.png)

**Figure 4** Forest plot of platelet reactivity at 1 hour postmorphine administration, using the VerifyNow method, subgroups according to study design. IV, inverse variance; RCTs, randomised controlled trials.
the risk of bias associated with this objective measure was considered to be low. Despite the widespread use of morphine in chest pain and anxiety relief in patients with ACS, conflicting data about its clinical impact has recently come to light. The activation of opioid receptors in the myenteric plexus decreases gut motility and secretion, inhibiting the activation of drugs whose action is directed at the P2Y12 protein and decreasing its absorption and bioavailability. Moreover, morphine is also known for its proemetic and antiperistaltic effects, which can further contribute to the decreased absorption of antiplatelet drugs.

This systematic review was planned and designed to evaluate the safety outcomes associated with morphine use in ACS. Pooled data RCTs and observational studies showed that treatment with morphine in patients with ACS is associated with a significant increase risk of in-hospital mortality, MACE and platelet reactivity. We found that morphine decreased the antiplatelet effect of P2Y12 inhibitors in the first hours of ACS. The clinical significance of this increase is uncertain, as the magnitude of this change is less than the difference between ticagrelor and clopidogrel in ACS, but appears to be at least twice as large as the impact of esomeprazole on the pharmacodynamics of clopidogrel. This effect of morphine ceases to be relevant at around the 8 hour mark. This may contribute to a delay in the onset of acute medical treatment, a greater prothrombotic milieu and more myocardial damage in patients with ACS. What is more, the analgesic effect of morphine followed by a decreased sympathetic response of the patient, without directly reversing the cause of ACS, may lead physicians to underestimate the severity of the underlying disease and to postpone the referral to an invasive revascularisation procedure. All the above-mentioned reasons may contribute towards the increased risk of in-hospital mortality and MACE related to the use of morphine. In clinical practice, other opioid analgesic drug such as fentanyl can be used, and a recent trial showed that fentanyl treatment in ACS increased platelet reactivity compared with no treatment. Although this suggests a possible class-effect of opioids on antiplatelet drugs, the evidence is sparse and requires further investigation before firm conclusions can be made.

Unexpectedly, we did not find an increased risk of nausea/emesis associated with morphine. This raises the likelihood that the reduction of gut secretion and motility is the core effect through which morphine decreases the activation of P2Y12 drugs.

With regard to platelet reactivity, we believe that the magnitude of the difference found supports a change in clinical practice, moving away from a recommendation to use morphine in ACS to recommending not using it routinely. The strength of this recommendation may be controversial due to the nature of the trials used and the other outcomes in this review not being statistically significant.

An important concern when combining randomised and observational data is the extent to which the participants and clinical setting are sufficiently similar to justify their pooling. On this account, the results of this review are robust since we found low heterogeneity across the outcomes of interest and the fact that none of the subgroup analyses comparing RCTs versus observational studies were statistically significant. Further proof of the consistency of the results is that no subgroup analysis showed a difference between STEMI and NSTEMI.

However, due to concerns over risk of bias across studies, we assessed the certainty in the evidence as low, despite there being little concern regarding inconsistency, indirectness or lack of statistical power.

**Figure 5** Forest plot of platelet reactivity at 2 hours postmorphine administration, using the VerifyNow method, subgroups according to study design. IV, inverse variance; RCTs, randomised controlled trials.
The key limitation of this review comes from the key limitation of most observational research, namely confounding. In a conservative approach, we attempted to minimise the impact of observational studies and their bias in the estimates by applying a correction factor previously used in other meta-analysis.26 27 Nevertheless, we must recognise that this adjustment is artificial and limits our results. Another limitation regards the possible differences in the doses and route of administration of morphine that were not available in most of the included studies.

Physicians may administer morphine to patients with more severe forms of chest pain, which may correspond to a more severe underlying ACS. This means that the increased risk of negative clinical outcomes could come as a result of patients being given morphine, or, alternatively, from the fact that morphine is usually reserved for the sickest patients. Because the included observational studies were substantially larger than the RCTs, including them in the meta-analysis could increase the risk of producing a biased result with an undue degree of statistical precision. To minimise this risk, we used methods to decrease the weight given to the largest and most biased studies, providing a more conservative estimate based on the available evidence. In doing so, we have produced the first and only systematic review to date that evaluates this highly relevant clinical question.

CONCLUSIONS

This systematic review raises concern about the use of morphine in patients with ACS and challenge the current clinical recommendations for its use in ACS. Most data come from studies at high risk of bias when evaluating the true effect of morphine in this setting. As such, a low-bias, adequately powered RCT designed to evaluate this question would be of significant scientific and clinical value. However, there is high certainty that morphine decreased the antiplatelet effect of P2Y12 inhibitors in the first hours of ACS.

Author affiliations
1Laboratory of Clinical Pharmacology and Therapeutics, Faculdade de Medicina, Universidade de Lisboa, Lisboa, Portugal
2Instituto de Medicina Molecular, Lisboa, Portugal
3Serviço de Cardiologia, Hospital Universitário de Santa Maria (CHLN), CAMIL, Centro Cardiovascular da Universidade de Lisboa - CCUL, Faculdade de Medicina, Universidade de Lisboa, Lisboa, Portugal
4Campus Neurológico Sénior, Torres Vedras, Portugal
5Laboratory of Clinical Pharmacology and Therapeutics, Universidade de Lisboa Faculdade de Medicina, Lisboa, Portugal
6Centro de Estudos de Medicina Baseada na Evidência, Universidade de Lisboa, Lisboa, Portugal

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Contributors
DC and GSD conceived the idea for the protocol and made the main contribution to planning and preparation of timelines for its completion. DC and GSD planned the data extraction and statistical analysis, as well as as risk of bias, quality of evidence and completeness of reporting assessments. GSD designed the tables and wrote the first draft of the manuscript, which was then reviewed and amended by ANF, DC, FBR, FP, JC and JF. All authors then approved the final written manuscript. DC is the guarantor for the work.

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JF received speaker and consultant fees from Abbott, Bial GlaxoSmithKline, Glunenthal, Lundbeck, Merck-Serono, Merck Sharp and Dohme, Novartis, TEVA, and Solvay. FJP received speaker and consultant fees from Astra Bayer, Boehringer Ingelheim, Daiichi and Sankyo Zeneca. All remaining authors declare no competing interests.

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No unpublished data are available.

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