# openheart Major bleeding after percutaneous coronary intervention and risk of subsequent mortality: a systematic review and meta-analysis

Chun Shing Kwok,<sup>1</sup> Sunil V Rao,<sup>2</sup> Phyo K Myint,<sup>3</sup> Bernard Keavney,<sup>1</sup> James Nolan,<sup>4</sup> Peter F Ludman,<sup>5</sup> Mark A de Belder,<sup>6</sup> Yoon K Loke,<sup>7</sup> Mamas A Mamas<sup>1</sup>

## ABSTRACT

**Objectives:** To examine the relationship between

periprocedural bleeding complications and major

outcomes following percutaneous coronary

meta-analysis of PCI studies that evaluated

assessed by considering the I<sup>2</sup> statistic.

intervention (PCI) and study differences in the

prognostic impact of different bleeding definitions.

Methods: We conducted a systematic review and

on MACEs and mortality outcomes. A systematic

search of MEDLINE and EMBASE was conducted to

identify relevant studies. Data from relevant studies

used to estimate the risk of adverse outcomes with

periprocedural bleeding. Statistical heterogeneity was

**Results:** 42 relevant studies were identified including

independently associated with increased risk of mortality

(OR 3.31 (2.86 to 3.82), I<sup>2</sup>=80%) and MACEs (OR 3.89

(3.26 to 4.64), I<sup>2</sup>=42%). A differential impact of major

bleeding as defined by different bleeding definitions on

REPLACE-2 (OR 6.69, 95% CI 2.26 to 19.81), STEEPLE

95% CI 1.74 to 16.74) had the worst prognostic impacts while HORIZONS-AMI (OR 1.51, 95% CI 1.11 to 2.05)

**Conclusions:** Major bleeding after PCI is independently

associated with a threefold increase in mortality and

MACEs outcomes. Different contemporary bleeding

outcomes, with 1.5-6.7-fold increases in mortality

observed depending on the definition of major bleeding

definitions have differential impacts on mortality

(OR 6.59, 95% CI 3.89 to 11.16) and BARC (OR 5.40,

533 333 patients. Meta-analysis demonstrated that

periprocedural major bleeding complications was

mortality outcomes was observed, in which the

had the least impact on mortality outcomes.

were extracted and random effects meta-analysis was

periprocedural bleeding complications and their impact

adverse cardiovascular events (MACEs) and mortality

**To cite:** Kwok CS, Rao SV, Myint PK, *et al.* Major bleeding after percutaneous coronary intervention and risk of subsequent mortality: a systematic review and meta-analysis. *Open Heart* 2014;**1**:e000021. doi:10.1136/openhrt-2013-000021

► Additional material is published online only. To view please visit the journal online (http://dx.doi.org/10. 1136/openhrt-2013-000021).

Received 13 December 2013 Revised 11 January 2014 Accepted 18 January 2014



For numbered affiliations see end of article.

#### **Correspondence to**

Dr Mamas A Mamas; mamasmamas1@yahoo.co.uk

# INTRODUCTION

used.

Advances in antithrombotic therapy have improved the outcomes of patients undergoing

# KEY MESSAGES

- The strength of this systematic review was the large number of studies included with over half a million total participants.
- Another strength was that we were able to evaluate the effect of different major bleeding definitions and its impact on risk of mortality and major adverse cardiovascular events.
- This systematic review had the limitation that studies included varied in antithrombotic and antiplatelet regimes after PCI procedure.
- Another limitation was that the systematic review was unable evaluate whether the subsequent mortality was directly related to the bleed.

percutaneous coronary intervention (PCI) through the reduction of ischaemic events at the expense of increased procedure-related bleeding complications. Major bleeding events in contemporary PCI are significant, with 30-day bleeding event rates reported between 0.7% and 1.1% in elective, <sup>1–3</sup> 0.6% and 4.7% <sup>3–6</sup> in non-ST-elevation myocardial infarction (NSTEMI) and 0.9% and 8.9% in ST-elevation myocardial infarction (STEMI) <sup>3 4 7 8</sup> depending on the definition used.

There are currently around 10 different definitions of major bleeding used in trials and registries of patients undergoing PCI9 10 and these definitions include various clinical events, such as blood transfusion or retroperitoneal haemorrhage, laboratory paraas differing values meters, such of haemoglobin decreases, and clinical out-comes such as mortality<sup>9</sup> resulting in significant differences in bleeding event recording across clinical trials thereby making comparisons between therapeutic strategies difficult. Furthermore, the incidence of major bleeding varies depending on definition used. In

one study, the RIVAL non-coronary artery bypass graft (non-CABG) related major bleeding occurred in 0.87% in the STEMI cohort and 0.57% in the NSTEMI group, while if an ACUITY major bleeding definition was used, major bleeding occurred in 3.1% in the STEMI group and 2.26% in the NSTEMI group, respectively.

Major periprocedural bleeding complications following PCI are predictors of mortality and major adverse cardiovascular events (MACEs),<sup>11–13</sup> with up to 12.1% of all in-hospital mortality after PCI in the National Cardiovascular Data Registry's CathPCI Registry related to bleeding complications.<sup>14</sup> In contrast, other studies have suggested that although bleeding may be causally related to adverse outcomes in some patients in the realworld setting, it is often merely a marker for patients at higher risk for adverse outcomes.<sup>15 16</sup>

Some prior studies that have reported on the prognostic impact of major bleeds have not accounted for differences in baseline covariates such as age, syndrome of presentation and comorbidities that would themselves impact on MACEs and mortality outcomes.<sup>17–21</sup> In contrast, while other studies have adjusted for baseline covariates, different definitions of major bleeding such as TIMI,<sup>2</sup> <sup>11</sup> <sup>22</sup> <sup>23</sup> GUSTO,<sup>20</sup> <sup>24</sup> STEEPLE<sup>20</sup> <sup>25</sup> and BARC<sup>23</sup> have been used, which have been shown to have differential impacts on mortality/MACEs outcomes.<sup>20</sup> <sup>23</sup> <sup>26</sup> Furthermore, the timing of bleeding from index PCI procedures included in such studies has varied from 48 h,<sup>25</sup> been limited to those that occur in hospital,<sup>23</sup> <sup>27</sup> to 30 days<sup>2</sup> <sup>28</sup> with impact on mortality and MACEs outcomes studied at different time points such as 30 days,<sup>8</sup> <sup>24</sup> <sup>28</sup> 6 months<sup>24</sup> <sup>27</sup> or 1 year.<sup>2</sup> <sup>22</sup> <sup>23</sup> <sup>25</sup>

Until today, there has not been a systematic review or meta-analysis previously published studying the prognostic impact of periprocedural major bleeding events on mortality and MACEs outcomes following PCI. We have therefore undertaken a meta-analysis to systematically study the impact of major bleeding following PCI on mortality and MACEs outcomes. In this meta-analysis, we provide an overview of the cohorts evaluating the rates of major bleeding events and systematically study the differences in the prognostic impact of different bleeding definitions and the relationship between major bleeding and clinical events at different time points.

## METHODS Eligibility criteria

We selected studies of patients who underwent PCI that reported on mortality or cardiovascular events among patients with and without major bleeding events. There was no restriction based on study design, definition of major bleeding or the indication for PCI or its status as an urgent or elective procedure. We excluded studies that did not report on categories of major bleeding and those that did not report either mortality or MACEs.

#### Search strategy

A search of EMBASE (1974 to January 2014) and MEDLINE (1946 to January 2014) was conducted on OvidSP. The search terms are shown in online supplementary figure S1. We did not use any language restrictions. We checked the bibliographies of included studies and relevant review articles found on the search for additional relevant articles.

#### Study selection and data extraction

Two reviewers (CSK and YKL) checked all titles and abstracts for studies that could potentially meet the inclusion criteria. We retrieved full reports of these potentially eligible studies and independently extracted data on study design, participant characteristics, interventions used, major bleeding events, follow-up, outcome events and methods of ascertaining measured clinical events on to a preformatted spreadsheet. Any discrepancies between the two reviewers were resolved by consensus after consulting a third reviewer (MAM).

#### Quality assessment

Risk of bias was assessed by considering four different areas: ascertainment of major bleeding events, ascertainment of outcome events, extent of loss to follow-up and the use of adjustment for confounders in the analysis. We also assess for publication bias using funnel plots when there were >10 studies available in the meta-analysis and there was no evidence of substantial statistical heterogeneity.<sup>29</sup>

#### Data analysis

We used RevMan V.5.1.7 (Nordic Cochrane Centre) to do random effects meta-analysis using the inverse variance methods for pooled ORs. We assumed similarity between the OR and other relative measures such as relative risk, rate ratios or HRs because cardiovascular events and death were rare events.<sup>30</sup> The analysis was stratified based on whether the results had considered the effect of potential confounders through adjustments or propensity-matched cohorts or not. In order to reduce the risk of bias from confounding so that we have a more reliable estimate of the independent effect of bleeding on prognosis, we appraised studies with multivariate adjustments or propensity-matched cohorts separately from studies with crude or unadjusted results. We used adjusted or propensity-matched risk estimates where available. For datasets reporting multiple time points, we took the earliest time point for the primary analysis. Any later time points were subsequently analysed in the separate Forest plots stratified according to timing of outcome assessment. Where there were multiple definitions of major bleeding we choose to use the results for TIMI major bleeding.

We planned for three analyses. The primary analysis was the pooled adjusted and unadjusted risk of mortality with and without major bleed. The secondary analysis was the unadjusted and adjusted risk of MACEs with and without major bleed. In a post hoc sensitivity analysis, we conducted a pooled analysis after excluding studies where it was clear that only some (and not all) of the participants had undergone PCI.

#### Statistical heterogeneity

We used the  $I^2$  statistic to assess statistical heterogeneity.  $I^2$  Values of 30–60% represent moderate levels of heterogeneity.<sup>31</sup>

#### RESULTS

## Study selection

The process of selection of studies is shown in online supplementary figure S1. We retrieved 42 relevant studies of patients that underwent PCI (total number of participants 533 333), which evaluated the risk of adverse events with and without major bleeding.<sup>2 8 11 14 17-23 27 28 32-58</sup> The number of participants in each study ranged from 352 to 280 390 and the number of major bleeding events was 66 277 (37 studies, 15.3%). A total of 40 studies evaluated mortality as an outcome and 11 studies reported on MACEs.

#### **Description of studies included**

The study designs, date of study, country of origin and indication for PCI is shown in table 1. There were 12 studies (149 650 participants), which were post hoc analyses of randomised controlled trials and 1 matched observational study (280 390 participants). Of the studies that reported number of centres, there were more multicentre studies than single centre studies (n=17 and n=13). The age and gender of participants along with the antiplatelet and anticoagulant regimes used is shown in online supplementary table S1.

Table 2 shows the timing of major bleeding, definition of major bleeding used in the individual studies and the incidence of major bleeding in each study. While many of the studies did not report when assessment for major bleeding took place (n=15), 15 of the studies clearly stated that they evaluated in-hospital major bleeding. The definitions of major bleeding also varied among the included studies and formal definitions of major bleeding included TIMI, GUSTO, STEEPLE, HORIZON-AMI, CRUSADE, BARC and REPLACE-2 definitions. These definitions are summarised in online supplementary table S2. Follow-up of patients in the studies included in this analysis varied from 48 h to more than 3 years. The impact of major bleeding on clinical outcomes was adjusted for baseline covariates in 22 studies (490 699 participants) while in 20 studies (42 634 participants) only unadjusted outcomes were reported.

## **Quality assessment**

Online supplementary table S3 shows the quality assessment for included studies. Ascertainment of bleeding

and mortality varied from data collection from medical record reviews to prospective evaluation in trials where independent committees adjudicated bleeding and outcome events.

Major bleeding and risk of mortality at any time point

The impact of major bleeding on mortality outcomes was studied in 40 studies reporting outcomes in 525 691 patients. In total, 66 016 major bleeds were reported. Crude rates or risk estimates for mortality in individual studies are shown in online supplementary table S4. Mortality rate was 3595/62 036 (5.8%, 28 studies) in patients who sustained a major bleed and 8937/370 522 (2.4%, 28 studies) in patients not experiencing major bleeding complications.

Meta-analysis of these data demonstrated that the overall risk of mortality was significantly greater among patients who sustained major bleeding complications periprocedurally (figure 1). The risk estimate for mortality was significantly lower in the subgroup of studies, which adjusted for potential confounders (OR 3.31 (2.86 to 3.82),  $I^2$ =80%, 491 565 participants compared to the OR calculated in those studies that did not adjust for baseline covariates; OR 6.75 (4.99 to 9.12),  $I^2$ =62%, 34 126 participants).

#### Major bleeding and risk of MACEs at any time point

The impact of major bleeding on MACEs outcomes was studied in 13 studies reporting outcomes in 69 843 patients. Crude rates or risk estimates for MACEs in individual studies are shown in tables 3 and 4; 757 major bleeds (6.8%) were reported.

MACE rates were 295/1701 (17.3%) in patients who sustained a major bleed and 2101/38 520 (5.4%) in patients not experiencing major bleeding complications. The overall risk of MACEs was significantly higher among patients with major bleeds (figure 2). The risk of MACEs did not significantly differ in the subgroup of studies that adjusted for baseline covariates (OR 3.89 (3.26 to 4.64),  $I^2$ =42%, 25 829 participants as compared to those that did not adjust for baseline covariates OR 3.12 (2.32 to 4.19),  $I^2$ =52%, 101 184 participants).

Mortality with major bleeding at different follow-up durations The adjusted risk of mortality was significantly greater among patients with major bleeds at all time points evaluated (figure 3). For 30-day mortality, adjusted risk was OR 3.24 (2.73 to 3.84, I<sup>2</sup>=81%, 11 studies, 403 457 participants) and at 6 months, the OR remained elevated at 3.23 (2.92 to 3.57, I<sup>2</sup>=0%, 4 studies, 50 872 participants; table 3). At the 1-year time point, the risk of mortality was OR 3.64 (2.39 to 5.56, I<sup>2</sup>=89%, 9 studies).

## MACEs with bleeding at different follow-up durations

Analysis of the impact of major bleeding on MACEs outcomes at different time points was stratified into risk of

Table 1 Study design	, year of study, country of	origin and participant inc	lusion criteria		
Study ID	Design	Date of study	Number of	Country	Inclusion criteria
				Obunity	
Amlani <i>et al</i>	Prospective cohort	May 2003 to July 2007	Single centre	Canada	Patients with STEMI
Ariza-Sole <i>et al</i> <sup>33</sup>	Prospective cohort	October 2009 to April 2012	Single centre	Spain	Patients with STEMI
Barthélémy et al 20	Prospective cohort	NA	Single centre	France	Patients in E-Paris Registry with STEMI and PCI
Bertrand et al 34	Post hoc analysis of RCT	October 2003 to April 2005	Single centre	Canada	Patients in EASY trial with PCI without STEMI
Boden <i>et al</i> <sup>35</sup>	Cohort	NA	NA	The Netherlands	Patients with STEMI and PCI
Budaj <i>et al</i> <sup>36</sup>	Post hoc analysis of RCT	NA	Multicentre	International	Patients with NSTEMI ACS. Note that only 64-79% had PCI
Cayla <i>et al</i> <sup>18</sup>	Post hoc analysis of RCT	NA	Multicentre	France	Patients in ABOARD trial with NSTEMI and PCI
Cayla <i>et al</i> 37	Post hoc analysis of RCT	NA	Multicentre	International	Patients in ATOLL study with STEMI and PCI
Chhatriwalla et al 14	Matched cohort	2004 to 2011	Multicentre	USA	Patients in the CathPCI Registry
Correia <i>et al</i> 38	Cohort	August 2007 to December 2010	Multicentre	Brazil	Patients with ACS. Note that only 76–90% had coronary angiogram
Eikelboom et al 28	Post hoc analysis of BCTs	NA	Multicentre	International	Patients in OASIS 1 and 2 and CURE who had ACS. Note that only 10–11% had PCI/stent/atherectomy
Fuchs <i>et al</i> <sup>21</sup>	Prospective cohort	January 2001 to June 2005	Single centre	Israel	Patients with STEMI and PCI
Gitt <i>et al</i> <sup>39</sup>	Cohort	October 2006 to October 2008	Multicentre	International	Patients with ACS. Note that only 65–82% had primary/rescue PCI
Giugliano <i>et al</i> <sup>8</sup>	Post hoc analysis of RCT	NA	Multicentre	International	Patients in ExTRACT-TIMI 25 trial where patients received fibrinolysis
Hermanides <i>et al</i> <sup>22</sup>	Prospective cohort	January 1991 to December 2004	Single centre	The Netherlands	Patients with STEMI and PCI
Kaul <i>et al</i> <sup>40</sup>	Post hoc analysis of RCT	May 2004 to August 2008	Multicentre	International	Patients with NSTEMI
Kikkert et al 41	Prospective cohort	January 2003 to July 2008	Single centre	The Netherlands	Patients with STEMI and PCI
Kinnaird et al 11	Retrospective cohort	1991 to 2000	NA	USA	Patients who underwent PCI
Le May et al 42	Cohort	May 2005 to July 2010	NA	Canada	Patients with STEMI and PCI
Lee et al 43	Prospective cohort	February 2003 to March 2006	NA	Korea	Patients with PCI with DES
Lemesle et al 27	Cohort	January 2000 to December 2007	Single centre	USA	Patients $\geq$ 80 years of age with PCI
Lindsey et al 44	Prospective cohort	July 2004 to January 2006	Multicentre	USA	Patients in EVENT registry with PCI
Lopes et al 45	Retrospective cohort	November 2001 to December 2006	Multicentre	USA	Patients in CRUSADES registry and subset with PCI

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## Table 1 Continued

			Number of		
Study ID	Design	Date of study	centres	Country	Inclusion criteria
Matic <i>et al</i> <sup>46</sup>	Cohort	August 2009 to December 2012	NA	Serbia	Patients with STEMI and PCI
Matic <i>et al</i> 47	Cohort	August 2009 to December 2010	NA	Serbia	Patients with STEMI and PCI
Matic <i>et al</i> <sup>48</sup>	Prospective cohort	August 2009 to December 2010	NA	Serbia	Patients with STEMI and PCI
Mehran <i>et al</i> <sup>2</sup>	Post hoc analysis of RCTs	NA	Multicentre	International	Patients in REPLACE-2, ACUITY and HORIZONS-AMI trial with PCI
Montalescot et al 25	Post hoc analysis of RCT	December 2005 to December 2006	Multicentre	International	Patients in STEEPLE trial who underwent elective PCI
Mrdovic <i>et al</i> <sup>49</sup>	Cohort	February 2006 to December 2009	Single centre	Serbia	Patients with STEMI and PCI
Musumeci <i>et al</i> <sup>50</sup>	Cohort	June 2005 to June 2008	Multicentre	Italy	Patients with PCI and DES
Ndrepepa <i>et al</i> <sup>17</sup>	Post hoc analysis of RCT	September 2005 to January 2008	NA	Germany	Patients in ISAR-REACT-3 trial with PCI
Ndrepepa <i>et al</i> <sup>23</sup>	Post hoc analysis of RCT	NA	NA	Germany	Patients in ISAR-REACT, ISAR-SWEET, ISAR-SMART and ISAR-REACT-2 trials with PCI
Pierre-Louis <i>et al</i> <sup>51</sup>	Cohort	NA	NA	USA	Patients with ACS and PCI
Pilgrim and Wenaweser 52	Cohort	May 2002 to December 2005	Single centre	Switzerland	Patients with PCI
Polanska-Skrzypczyk <i>et al</i> <sup>53</sup>	Prospective cohort	February 2001 to October 2002	Single centre	Poland	Patients with STEMI and PCI
Poludasu <i>et al</i> <sup>54</sup>	Prospective cohort	July 2001 to May 2010	Multicentre	USA	Patient with PCI
Rao <i>et al</i> <sup>24</sup>	Post hoc analysis of RCTs	NA	Multicentre	International	Patients in GUSTO IIb, PURSUIT and PARAGON A/B. Note that only 11–30% had PCI
Rossini <i>et al</i> <sup>55</sup>	Cohort	NA	NA	USA	Patients with PCI and DES
Urban <i>et al</i> <sup>56</sup>	Prospective cohort	May 2006 to April 2008	Multicentre	International	Patients in e-SELECT registry with PCI and DES
Valente et al 19	Prospective cohort	January 2004 to December 2008	Single centre	Italy	Patients in Intensive Cardiac Care Florence STEMI Registry who had STEMI and PCI
Yoon <i>et al</i> 57	Cohort	NA	NA	Korea	Patients in IRS DES registry with PCI and DES
Zheng <i>et al</i> <sup>58</sup>	Retrospective cohort	January 2004 to January 2008	Single centre	China	Patients with elective or urgent PCI

ACS, acute coronary syndromes; DES, drug-eluting stent; NA, not applicable; NSTEMI, non-ST-elevation myocardial infarction; PCI, percutaneous coronary intervention; RCT, randomised controlled trial; STEMI, ST-elevation myocardial infarction.

Table 2	Timing of bleeding,	definition of major	bleeding and follow-up
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Chudu ID	Timing of		Number of participants in	Number of participants in	<b>Fallow</b>
Study ID	bleeding	Major/severe bleeding criteria	bleeding group	control group	Follow-up
Amlani <i>et al</i> <sup>32</sup>	Within 30 days	Haemoglobin drop $\geq$ 5 g/dL, intracranial haemorrhage, bleeding requiring surgery or blood transfusion of at least	152 (67 had PCI)	1237 (566 had PCI)	30 days
Ariza-Sole et al 33	In-hospital	CRUSADE	33	1031	Mean follow-up 344 days
Barthélémy <i>et al</i> <sup>20</sup> Bertrand <i>et al</i> <sup>34</sup>	In-hospital Unclear	TIMI, GUSTO, STEEPLE REPLACE-2	Total 671 19	NA 1329	1 year 30 days, 6 months, 1 year
Boden <i>et al</i> <sup>35</sup> Budaj <i>et al</i> <sup>36</sup>	In-hospital Up to 180 days	CRUSADE ESSENCE	203 771 (30 days), 937	762 18 851 (30 days),	1 year 180 days
Cayla <i>et al</i> 18	30 days	STEEPLE	(180 days) 19	18 851(180 days) 333	30 days
Cayla <i>et al</i> <sup>37</sup> Chhatriwalla <i>et al</i> <sup>14</sup>	30 days In-hospital	STEEPLE CathPCI Registry definition	42 56 078	868 224 312	30 days In-hospital
Correia <i>et al</i> $^{38}$ Eikolboom <i>et al</i> $^{28}$	In-hospital Within 20 days	Blooding that was significantly disabling blooding requiring	29 792	426	In-hospital
	Within 30 days	transfusion of $\geq 2$ units of packed cells or bleeding that was life threatening	700	3303	SO-day monanty
Fuchs <i>et al</i> <sup>21</sup>	Unclear	TIMI	27	804	30 days and 6 months
Gitt et al 39	In-hospital	Drop in haemoglobin of >5 g/dL or haematocrit of >15%	281	8451	In-hospital
Giugliano <i>et al</i> <sup>8</sup>	Up to day 8	ТІМІ	309	20 014	30 days
Hermanides et al 22	48 h	TIMI	80	4371	30 days and 1 year
Kaul <i>et al</i> <sup>40</sup>	Up to 120 h after randomisation	GUSTO	598	8808	30 days
Kikkert et al 41	Unclear	ТІМІ	35	331	1–30 days
Kinnaird <i>et al</i> <sup>11</sup>	Unclear	ТІМІ	588	8992	In-hospital
Le May et al 42	Unclear	ТІМІ	91	1941	6 months
Lee et al 43	Median 1366 days	STEEPLE	148	3022	Median 1366 days
Lemesle <i>et al</i> <sup>27</sup>	In-hospital	Decrease in haematocrit of $\geq$ 15% and/or the occurrence of a major haematoma/gastrointestinal bleeding/intracerebral bleeding	127	2639	6 months
Lindsey et al 44	In-hospital	TIMI	180	5781	1 year
Lopes et al 45	In-hospital	CRUSADE	3902	28 993	30 days, 1 year, 3 year, >3 years
Matic et al 46	Unclear	GUSTO	32	738	In-hospital
Matic et al 47	Unclear	HORIZONS-AMI	88	1154	30 day
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Study ID	Timing of bleeding	Major/severe bleeding criteria	Number of participants in bleeding group	Number of participants in control group	Follow-up
Matic <i>et al</i> 48	Unclear	BARC	114	1558	1 year
Mehran et al 2	30 days	TIMI	267	17 034	Up to 1 year
Montalescot et al 25	48 h	STEEPLE	3528 (total)	NA	1 year
Mrdovic <i>et al</i> <sup>49</sup>	Unclear	TIMI	22	2074	30 days and 1 vear
Musumeci <i>et al</i> 50	30 days, 12 months, any follow-up	TIMI	52	1385	3 years
Ndrepepa et al 17	Unclear	REPLACE-2	555	4015	1 year
Ndrepepa et al 23	In-hospital	TIMI, REPLACE-2, BARC	Total 12 459	NA	30 days, 1 year
Pierre-Louis <i>et al</i> <sup>51</sup>	In-hospital	Intracerebral or intraocular bleeding, clinical bleeding requiring blood transfusion, clinical bleeding with a reduction in haematocrit >10 points, retroperitoneal or gastrointestinal bleeding, access site bleeding requiring intervention, and $\geq$ 4 cm diameter vascular access site haematoma	34	600	In-hospital
Pilgrim and Wenaweser <sup>52</sup>	In-hospital	TIMI	48	3787	30 days
Polanska-Skrzypczyk et al 53	In-hospital	TIMI	40	1024	1 year
Poludasu et al 54	Unclear	HORIZONS-AMI	396	11 595	2.3 years
Rao et al 24	Unclear	GUSTO	2908	23 544	Up to 6 months
Rossini <i>et al</i> 55	Unclear	TIMI	57	1301	1 year
Urban <i>et al</i> 56	Unclear	STEEPLE	Total 15 147	NA	360 days
Valente et al 19	In-hospital	TIMI, ACUITY	Total 991	NA	In-ICCU
Yoon <i>et al</i> 57	In-hospital	BARC type 2–5	234	5932	2 years
Zheng et al 58	Unclear	TIMI	27	385	1 year

				Odds Ratio	Odds Ratio
Study or Subgroup lo	og[Odds Ratio]	SE	Weight	IV, Random, 95% CI	IV, Random, 95% CI
1.1.1 Unadjusted analys	sis				
Ariza-Sole 2013	1,93297	0.315633	2.4%	6.91 [3.72, 12.83]	
Barthelemev 2012	2.952825	0.521713	1.4%	19.16 (6.89, 53.27)	
Bertrand 2009	5.939803	1.567807	0.2%	379.86 [17.58.8206.22]	
Boden 2012	1.396245	0.334086	2.3%	4.04 [2.10, 7.78]	
Cavla 2011	3 918005	0.817273	0.7%	50 30 (10 14 249 59)	
Cavla 2012	1 871802	0.421276	1.8%	6 50 [2 85 14 84]	
Fuchs 2009	1 449269	0.459527	1.6%	4 26 [1 73 10 48]	
Kinnaird 2003	1 508512	0.174428	3.6%	4 52 [3 21 6 36]	
Lee 2009	1 818077	0 274042	2.7%	6 16 [3 60 10 54]	
Matic 2010	1 261 298	0.644726	1.0%	3 53 [1 00 12 49]	
Matic 2011	0 746688	0.357863	2.2%	2 11 [1 05 4 25]	
Mrdovic 2013	2 701 361	0.446599	17%	14 90 (6 21 35 75)	
Musumeri 2012	1 353255	0.385864	2.0%	3 87 [1 87 8 74]	
Ndrenena 2010	1 798404	0.320035	2.0%	6 04 [3 23 11 31]	
Pierre 2010	4 864298	1 489577	0.2%	129 58 6 99 2401 421	
Pilarim 2010	2 200552	0.456042	1.6%	9 03 [3 69 22 07:42]	
Rossini 2010	2.200332	0.430042	7.0%	7 55 [3 87 14 97]	
Volente 2011	2.021040	0.347304	2.2.0	9 34 [4 39 16 99]	
Subtotal (95% CI)	2.121005	0.320733	32.4%	6.75 [4.99, 9.12]	•
Heterogeneity: Tou <sup>2</sup> – 0.1	22: Chiz - 11 66	df = 17 (P	- 0.0003	· IZ - 62%	•
Telefoyenelly, rau = 0	23, CIII - 44.00 - 12 12 /P - 0.00	, ui = 17 (F 2004)	- 0.0003)	,1 = 02%	
restion overall ellect. Z -	- 12.43 (F < 0.00	5001)			
1.1.2 Adjusted analysis					
Amlani 2010	1.131402	0.357863	2.2%	3.10 [1.54, 6.25]	
Budaj 2009	1.241269	0.145547	3.8%	3.46 [2.60, 4.60]	-
Chhatriwalla 2013	1.068153	0.024481	4.4%	2.91 [2.77, 3.05]	•
Correia 2012	1.205971	0.527799	1.3%	3.34 (1.19, 9.40)	
Eikelboom 2006	1.680828	0.153983	3.7%	5.37 [3.97, 7.26]	-
Gitt 2010	0.779325	0.209011	3.3%	2.18 (1.45, 3.28)	
Giugliano 2010	1.064711	0.103435	4.1%	2.90 [2.37, 3.55]	+
Hermanides 2010	1.252763	0.217727	3.2%	3.50 (2.28, 5.36)	
Kaul 2013	1.388791	0.17187	3.6%	4.01 [2.86, 5.62]	
Kikkert 2013	1.007958	0.258763	2.9%	2 74 [1 65 4 55]	
Le May 2011	1 497388	0.359071	21%	4 47 [2 21 9 04]	
Lemesle 2009	1 098612	0.212477	3.2%	3 00 [1 98 4 55]	
Lindsev 2009	1 924249	0.36129	21%	6 85 [3 37 13 91]	
Lones 2012	0.693147	0.00120	3.6%	2 00 [1 44 2 77]	
Matic 2012	1.098612	0.293016	2.6%	3 00 [1 69 5 33]	
Mehran 2011	1 435085	0.155373	3.7%	4 20 [3 10 5 20]	
Montalescot 2009	1.098612	0.518602	1 4 %	3 00 [1 09 8 29]	
Ndrenena 2012	1 144223	0.510002	3.7%	3 14 [2 30 4 29]	
Polaneka 2012	1 788421	0.133023	2.0%	5 98 12 78 12 841	
Poludacu 2011	0.336472	0.0000000	2.0%	1 /0 /1 06 1 9/1	
Pag 2006	1 474762	0.13334	1 206	1.40 [1.00, 1.04]	+
Nau 2003 Urban 2011	1.474703	0.074302	9.370	4.37 (3.77, 3.00) 6 00 (3.22, 10.06)	
Subtotal (95% CI)	1.751735	0.302433	67.6%	3.31 [2.86, 3.92]	•
Hotorogonoity: Tou? - 0.1	07· Chiz - 100 7	0 df= 24 //	01.0/0	11)· IZ = 00%	•
Test for overall effect: Z =	= 16.32 (P < 0.0)	0, ui - 21 (r 3001)	~ 0.0000	51),1 = 60%	
T-4-1 (050) ON			100.00		
1 otal (95% CI)			100.0%	4.14 [3.59, 4.78]	
Heterogeneity: Tau <sup>2</sup> = 0.1	12; Chi² = 203.2	7, df = 39 (F	P < 0.0000	01); I² = 81%	
Test for overall effect: Z =	= 19.44 (P < 0.00	0001)			Favours major bleed Favours no major bleed
Test for subgroup differe	ences: Chi <sup>2</sup> = 17	55 df = 1 (	P < 0 000	1) I <sup>2</sup> = 94.3%	



MACE <1 year and  $\geq 1$  year due to the limited number of studies presenting MACE data and its relationship with time. The adjusted risk of MACE <1 year was OR 3.96 (3.26 to 4.81, I<sup>2</sup>=53%, 2 studies, 47 422 participants) and for  $\geq 1$  year was OR 3.19 (1.89 to 5.37, 1 study, 3170 participants; table 3).

Adverse outcomes and different definitions for major bleeding

The impact of different definitions of major bleeding on mortality and MACEs outcomes are presented in table 4. REPLACE-2 (OR 6.69 (2.26 to 19.81),  $I^2$ =84%, 3 studies, 17 996 participants) definition of major bleeding had

 Table 3
 Summary of risk of mortality and MACEs among patients with and without major bleed after percutaneous coronary intervention at different time points

Duration	OR	l² (%)	Number of participants	Number of events in bleed group/total	Number of events in non-bleed group/total				
Adjusted risk of mortality									
30 days <sup>8</sup> <sup>14</sup> <sup>24</sup> <sup>28</sup> <sup>32</sup> <sup>36</sup> <sup>38</sup> <sup>39</sup> <sup>40</sup> <sup>41</sup> <sup>45</sup>	3.24 (2.73 to 3.84)	81	403 457	3307/59 630	7645/317 375				
6 months <sup>24</sup> <sup>27</sup> <sup>36</sup> <sup>42</sup>	3.23 (2.92 to 3.57)	0	50 872	154/1028	1119/20 606				
1 year <sup>2 22 23 25 44 48 53 56</sup>	3.64 (2.39 to 5.56)	89	74 637	160/2494	677/36 167				
Adjusted risk of MACEs									
<1 year <sup>24 34 36</sup>	3.96 (3.26 to 4.81)	53	47 422	175/790	1204/20 180				
>1 year <sup>43</sup>	3.19 (1.89 to 5.38)	NA	3170	NA	NA				
MACEs, major adverse cardiovascular eve	ents; NA, not applicable	MACEs, major adverse cardiovascular events; NA, not applicable.							

intervention with different definitions of bleeding									
Mortality				MACEs					
Definition of major bleed	OR	l <sup>2</sup> (%)	Number of participants	Definition	OR	l <sup>2</sup>	Number of participants		
BARC <sup>23 38 48</sup>	5.40 (1.74 to 16.74)	88	14 550	BARC	NA	NA	NA		
CRUSADE <sup>33 35 45</sup>	3.69 (1.68 to 8.12)	85	14 055	CRUSADE	NA	NA	NA		
GUSTO <sup>24 40 46</sup>	4.30 (3.76 to 4.92)	0	27 222	GUSTO <sup>24</sup>	4.37 (3.78 to 5.07)	NA	26 452		
HORIZON-AMI <sup>47 54</sup>	1.51 (1.11 to 2.05)	12	13 233	HORIZON-AMI	NA	NA	NA		
REPLACE <sup>17 23 34</sup>	6.69 (2.26 to 19.81)	84	17 996	REPLACE	NA	NA	NA		
STEEPLE <sup>18 25 37 43 56</sup>	6.59 (3.89 to 11.16)	53	23 107	STEEPLE <sup>43</sup>	3.19 (1.89 to 5.37)	NA	3170		
TIMI <sup>2 8 11 19 20 21 22 42</sup> 44 49 50 52 53 55	5.15 (4.01 to 6.61)	69	69 449	TIMI <sup>21</sup>	2.41 (1.42 to 4.35)	NA	1348		
Other <sup>14 27 28 32 36 39 51</sup>	3.34 (2.59 to 4.32)	76	346 670	Other <sup>36</sup>	3.99 (3.3 to 4.82)	NA	19 622		
MACEs, major adverse cardiov	ascular events; NA, not	applicat	ole.						

 Table 4
 Summary of risk of mortality and MACEs among patients with and without major bleed after percutaneous coronary intervention with different definitions of bleeding

the greatest impact on mortality outcomes, while the ACUITY/HORIZONS-AMI had the least impact (OR 1.51 (1.11 to 2.05),  $I^2$ =12%, 1 study, 13 233 participants).

#### Sensitivity analysis

Sensitivity analysis excluding studies where not all participants had PCI did not significantly alter the pooled analysis for association between major bleeding and mortality (OR 3.16 (2.61 to 3.81),  $I^2=75\%$ , 372 449 participants).

#### DISCUSSION

Advances in antithrombotic and antiplatelet therapy have improved outcomes following PCI through the reduction of ischaemic events although this has been at the expense of increased procedure-related bleeding complications. Our meta-analysis of 42 studies including over half a million patients confirms that major bleeding is independently associated with a threefold increase in mortality and MACEs outcomes, and that this increased mortality and MACEs risk observed following a major bleed is sustained for periods of over 1 year. We also show that major bleeding events as defined by different contemporary bleeding definitions have differential impacts on mortality outcomes, with 1.5–6.7-fold increases in mortality observed depending on the definition of major bleeding used.

There are a number of potential mechanisms that may underlie the relationship between major bleeding and adverse mortality outcomes. Patients that sustain major bleeding complications post PCI are more likely to be older, have renal failure, undergo PCI for STEMI/ NSTEMI presentations, present with haemodynamic compromise or have a history of heart failure<sup>2 59</sup> that also

				Odds Ratio	Odds Ratio		
Study or Subgroup	log[Odds Ratio]	SE	Weight	IV, Random, 95% CI	IV, Random, 95% CI		
2.1.1 Unadjusted ana	lysis						
Cayla 2011	1.050822	0.550383	2.7%	2.86 [0.97, 8.41]			
Fuchs 2009	1.085189	0.43562	4.0%	2.96 [1.26, 6.95]	<b>.</b>		
Hermanides 2010	0.307485	0.596932	2.4%	1.36 [0.42, 4.38]			
Kinnaird 2003	1.150572	0.181399	10.9%	3.16 [2.21, 4.51]			
Matic 2011	1.199965	0.287357	7.0%	3.32 [1.89, 5.83]	_ <b></b>		
Musumeci 2012	1.345472	0.295607	6.8%	3.84 [2.15, 6.85]			
Rossini 2010	1.83737	0.278382	7.3%	6.28 [3.64, 10.84]			
Yoon 2013	0.647103	0.173044	11.2%	1.91 [1.36, 2.68]			
Zheng 2009	1.321756	0.541276	2.8%	3.75 [1.30, 10.83]			
Subtotal (95% CI)			55.1%	3.12 [2.32, 4.19]	•		
Heterogeneity: Tau <sup>2</sup> =	0.09; Chi <sup>2</sup> = 16.67	, df = 8 (P =	0.03); l² =	= 52%			
Test for overall effect:	Z = 7.54 (P < 0.00)	001)					
2.1.2 Adjusted analys	sis						
Bertrand 2009	0.879627	0.285592	7.1%	2.41 [1.38, 4.22]	_ <b></b>		
Budaj 2009	1.383791	0.096646	14.7%	3.99 [3.30, 4.82]	-		
Lee 2009	1.160021	0.266391	7.7%	3.19 [1.89, 5.38]			
Rao 2005	1.474763	0.074902	15.5%	4.37 [3.77, 5.06]			
Subtotal (95% CI)			44.9%	3.89 [3.26, 4.64]	♦		
Heterogeneity: Tau <sup>2</sup> =	0.01; Chi <sup>2</sup> = 5.15,	df = 3 (P = 0	).16); I <sup>2</sup> =	42%			
Test for overall effect:	Z = 15.12 (P < 0.0)	0001)					
Total (95% CI)			100.0%	3.34 [2.75, 4.06]	•		
Heterogeneity: Tau <sup>2</sup> =	0.06; Chi <sup>2</sup> = 30.48	, df = 12 (P	= 0.002);	I <sup>2</sup> = 61%			
Test for overall effect:	Z = 12.14 (P < 0.0)		Eavours major bleed Eavours no major bleed				
Test for subgroup differences: Chi <sup>2</sup> = 1.61, df = 1 (P = 0.20), l <sup>2</sup> = 38.0%							

Figure 2 Adjusted risk of major adverse cardiovascular event with and without major bleed after percutaneous coronary intervention.



**Figure 3** Risk of mortality with major bleed with different duration of follow-up.

independently predict adverse mortality outcomes themselves. Many previous studies<sup>17-21</sup> that have reported on the prognostic impact of major bleeds have not accounted for differences in such baseline covariates that are themselves known to impact on mortality outcomes, which would overestimate the mortality risks associated with a major bleed. For example, in an analysis from the Global Registry of Acute Coronary Events (GRACE), major bleeding was no longer associated with 6-month mortality after adjustment for the known comorbidities with the authors concluding that comorbidities associated with major bleeding accounting for the higher rate of mortality in patients who bled.<sup>15</sup> Consistent with this, we have observed that in meta-analysis of studies that do not adjust for baseline covariates, major bleeding is associated with a sixfold increased risk in mortality, but this decreases to a threefold independent increase in mortality once baseline covariates are adjusted for, suggesting that major bleeding is independently linked to mortality and MACEs outcomes.

The potential mechanisms by which a major bleed adversely impacts on clinical outcomes are numerous. Major bleeds such as an intracranial haemorrhage or severe blood loss as can occur in a gastrointestinal haemorrhage may result in mortality directly, although such bleeding events would not explain the persistent mortality risk observed for over a year following a major bleeding event in our meta-analysis. Major bleeds may necessitate the discontinuation of antiplatelet or antithrombotic medications that increase the risk for stent thrombosis, a strong independent predictor of mortality outcomes in the short and longer term (>1 year).60 Furthermore, increased production of erythropoietin in response to anaemia that occurs following a major bleed may contribute to a prothrombotic systemic state beyond the acute phase through platelet activation and induction of plasminogen activator inhibitor-1 (PAI-1).61 62 Treatment with erythropoietin has been associated with increased risk of thrombosis in critical care patients<sup>63</sup> and increase in the composite endpoint of death, acute myocardial infarction, stroke and stent thrombosis in patients with STEMI.<sup>64</sup> Blood transfusions have been linked to adverse shorter and longer term mortality<sup>65</sup> and have been shown to predict 30-day mortality<sup>66</sup> independently of bleeding and haematocrit. Adverse mortality outcomes associated with blood transfusions occur through a number of mechanisms including prothrombotic effects mediated through acute platelet release of CD40 ligand,<sup>67</sup> platelet activation and induction of PAI-1<sup>61</sup> an inhibitor of endogenous fibrinolytic mechanisms. These increased risks of mortality also extend to non-red blood cell transfusions such as platelets or plasma/cryoprecipitate that may also be utilised following a periprocedural major bleed.<sup>68</sup>

Our observations of a differential impact of different major bleeding definitions on mortality outcomes is particularly pertinent suggesting that the choice of bleeding definition used has significantly influenced the outcome of previous studies, since the definition employed will influence the prevalence of reported bleeds as well as their prognostic impact. For example, in the RIVAL study<sup>69</sup> non-CABG related major bleeding as defined by the study was not significantly different between the radial and femoral arms of the study (OR=0.73, 95% CI 0.43 to 1.23; p=0.9), while use of the ACUITY major bleeding criterion was associated with a statistically significant reduction in major bleeding in the radial arm of the study (OR=0.43, 95% CI 0.32 to 0.57; p<00001). Similarly, in the SYNERGY trial that used the GUSTO and TIMI major definition of bleeding, the enoxaparin arm was associated with a significantly higher rate of major bleeding as defined by TIMI major criteria but no difference in major bleeding as defined by the GUSTO major definition. Definitions that encompass less severe bleeding events (such as the ACUITY definition) are not as strongly linked to adverse events and therefore may not be a powerful means of evaluating bleeding avoidance strategies. In order to prove efficacy for bleeding avoidance the use of a more discriminative definition would be preferable.

Our meta-analysis has a number of potential limitations. First, an inherent limitation of any meta-analysis is that of publication bias; studies that show a neutral outcome in mortality are less likely to be published than those that show a positive outcome and thus tend to bias any meta-analysis of published data towards a more positive outcome. Second, studies included in this meta-analysis often used different antithrombotic and antiplatelet regimes for the PCI procedures undertaken for different indications, hence it is unclear whether the prognostic impact of a major bleed differs with different antiplatelet/anticoagulant combinations or whether it differs in the elective/acute coronary syndrome setting. Finally, our current analysis does not provide insight into whether the timing of the major bleed in relation to the index PCI procedure has a differential impact on mortality outcomes.

In conclusion, our meta-analysis of 42 studies including over half a million patients has revealed that major bleeding is independently associated with a threefold increase in mortality and MACEs outcomes, and that this increased mortality and MACEs risk observed following a major bleed is observed for periods of over 1 year. We also show that major bleeding events as defined by different contemporary bleeding definitions have differential impacts on mortality outcomes. Given the significant impact of major bleeding on mortality outcomes, formal bleeding risk assessment should be undertaken as part of the decision-making process for PCI procedures and bleeding avoidance strategies such as optimal pharmacotherapy and access site choice should be actively undertaken, particularly in those patients at highest baseline risk for bleeding complications.

#### Author affiliations

<sup>1</sup>Cardiovascular Institute, University of Manchester, Manchester, UK <sup>2</sup>Department of Cardiology, Duke Clinical Research Institute, Duke University

Medical Center, Durham, North Carolina, USA

<sup>3</sup>Division of Applied Health Sciences, School of Medicine & Dentistry, University of Aberdeen, Aberdeen, Scotland, UK

<sup>4</sup>Department of Cardiology, University Hospital North Staffordshire, Stoke-on-Trent, UK

<sup>5</sup>Department of Cardiology, Queen Elizabeth Hospital, Birmingham, UK
<sup>6</sup>Cardiothoracic Division, The James Cook University Hospital, Middlesbrough, UK

<sup>7</sup>Norwich Medical School, University of East Anglia, Norwich, UK

**Contributors** CSK, MAM and YKL contributed to planning, conduct and reporting of the work. SVR, PKM, BK, JN, PFL and MDB contributed to the interpretation of the findings and reporting of the work. MAM is the guarantor and was responsible for the overall content.

Competing interests None.

Provenance and peer review Not commissioned; externally peer reviewed.

Data sharing statement No additional data are available.

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