Diagnostic features, management and prognosis of type 2 myocardial infarction compared to type 1 myocardial infarction: a systematic review and meta-analysis

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ABSTRACT

Importance Distinguishing type 2 (T2MI) from type 1 myocardial infarction (T1MI) in clinical practice can be difficult, and the management and prognosis for T2MI remain uncertain.

Objective To compare precipitating factors, risk factors, investigations, management and outcomes for T2MI and T1MI.

Data sources Medline and Embase databases as well as reference list of recent articles were searched January 2009 to December 2020 for term ‘type 2 myocardial infarction’.

Study selection Studies were included if they used a universal definition of MI and reported quantitative data on at least one variable of interest.

Data extraction and synthesis Data were pooled using random-effect meta-analysis. Risk of bias was assessed using Newcastle-Ottawa quality assessment tool. Preferred Reporting Items for Systematic Reviews and Meta-analyses guidelines were followed. All review stages were conducted by two reviewers.

Main outcomes and measures Risk factors, presenting symptoms, cardiac investigations such as troponin and angiogram, management and outcomes such as mortality.

Results 40 cohort studies comprising 98,930 patients with T1MI and 13,803 patients with T2MI were included. Compared with T1MI, patients with T2MI were: more likely to have pre-existing chronic kidney disease (OR 1.87; 95% CI 1.53 to 2.28) and chronic heart failure (OR 2.35; 95% CI 1.82 to 3.03), less likely to present with typical cardiac symptoms of chest pain (OR 0.19; 95% CI 0.13 to 0.26) and more likely to present with dyspnoea (OR 2.64; 95% CI 1.86 to 3.74); more likely to demonstrate non-specific ST-T wave changes on ECG (OR 2.62; 95% CI 1.81 to 3.79) and less likely to show ST elevation (OR 0.22; 95% CI 0.17 to 0.28); less likely to undergo coronary angiography (OR 0.09; 95% CI 0.06 to 0.12) and percutaneous coronary intervention (OR 0.06; 95% CI 0.04 to 0.10) or receive cardioprotective medications, such as statins (OR 0.25; 95% CI 0.16 to 0.38) and beta-blockers (OR 0.45; 95% CI 0.33 to 0.63). T2MI had greater risk of all cause 1-year mortality (OR 3.11; 95% CI 1.91 to 5.08), with no differences in short-term mortality (OR 1.34; 95% CI 0.63 to 2.85).

Strengths and limitations of this study

- Inclusion of all contemporary cohort studies in the troponin era.
- Analyses of large population of patients with type 2 myocardial infarction and type 1 myocardial infarction which provided high level of precision.
- Wide array of clinically significant variables assessed providing a comprehensive analysis.
- Analysis of crude mortality only was possible due to lack of individual patient data.

Conclusion and relevance This review has identified clinical, management and survival differences between T2MI and T1MI with greater precision and scope than previously reported. Differential use of coronary revascularisation and cardioprotective medications highlight ongoing uncertainty of their utility in T2MI compared with T1MI.

INTRODUCTION

The clinical definition of myocardial infarction (MI) has evolved over time. The 2007 Universal Definition of Myocardial Infarction included a subset of MI that was secondary to aetiologies unrelated to underlying occlusive coronary artery disease.1 In 2012, the Third Universal Definition of Myocardial Infarction Consensus Document2 gave rise to the aetiologic distinction between T1MI, defined as MI due to plaque erosion and/or rupture, and T2MI, defined as MI caused by increased oxygen demand or decreased blood supply, in the absence of acute plaque rupture or coronary thrombosis. More recently, in 2018, the Fourth Universal definition of MI updated concepts of T2MI regarding specific situations associated with oxygen demand and supply imbalance and the relevance of the presence or
absence of underlying coronary artery disease to therapy and prognosis. (see online supplemental table S1 for more detail).

In clinical practice, distinguishing T2MI from T1MI based on clinical presentation, ECG features and cardiac troponin (cTn) values can be difficult. In the absence of randomised controlled trials that have evaluated different investigational and therapeutic interventions in patients with T2MI, uncertainty remains around the appropriate management of such patients, particularly those with known or suspected coronary artery disease. Past reviews have assessed one or more attributes of T2MI in comparison to T1MI but, to our knowledge, none have undertaken a comprehensive analysis of symptoms, physical signs, investigation results, management regimens and clinical outcomes, both short and long term, of T2MI versus T1MI.

We undertook a systematic review of observational studies with the aims of identifying diagnostic and investigational findings which can assist clinicians to better distinguish T2MI from T1MI, and compare T2MI with T1MI in defining differences in management strategies and clinical outcomes.

METHODS
Study design
The review was undertaken in accordance with recommendations of the Cochrane Collaboration and Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) statement. Our review was registered on PROSPERO prior to commencement (registration number: CRD42021237746). Medline and Embase databases were searched for all studies published between 1 January 2009, and 31 December 2020, using search terms to identify all studies related to T2MI (see online supplemental table S2). Reference lists of all relevant articles were also assessed to identify additional relevant studies. The study PRISMA flow chart is shown in online supplemental figure S1. January 2009 was chosen as the start date for the literature search in order to restrict our analyses to contemporary studies in the troponin era that employed formal definitions of T2MI which were only devised from 2007 onwards.

Studies were included if they: (1) compared patient populations with T2MI and T1MI, (2) used a universal definition of MI, (3) included at least one variable of interest, (4) were available as full text in English and (5) were either a randomised control trial or comparative observational study. Studies were excluded if: (1) no full text was available, (2) duplicate data were used or (3) less than 200 participants in total were included. Initial screening of titles and abstracts for eligible studies was performed independently by two authors (MK, KW), as was full-text review for inclusion, with any differences in review settled by consensus agreement.

Data collection and synthesis
Data pertaining to all variables of interest were collected from all included studies using a standardised pro forma by one author (MK) and independently reviewed by the second author (KW). These variables comprised: study dates, design, sample size, definition used to define T2MI and T1MI, patient demographics, pre-existing medical conditions, precipitating factors, clinical symptoms, ECG findings, laboratory values, echocardiographic results, any clinical interventions or medical treatments administered, and clinical outcomes observed.

Data on variables reported as, or able to be converted to, raw numbers, were pooled from all studies and subject to comparative meta-analysis using Review Manager (RevMan, Computer program. Version 5.3. Copenhagen: The Nordic Cochrane Centre, The Cochrane Collaboration, 2014). For each variable, the odds ratio (OR) comparing T2MI to T1MI, and its 95% confidence interval (CI), was calculated and weighted using the random effects method. As specified in the registered study protocol, the random effects method was used in anticipation of study heterogeneity of at least moderate degree (I² statistic of heterogeneity >50%). In addition to the weighted OR, we also report the crude total event rates for each variable subject to meta-analysis in order to provide a more clinically meaningful estimate of the prevalence of these events in each patient group in view of the large sample sizes. Studies reporting mean or median values only were reproduced as reported in the original study.

Risk of bias within each study was assessed using the Newcastle-Ottawa quality assessment tool for cohort studies, with scores 7–8 denoting good quality studies, 4–6 fair quality, and 0–3 poor quality. Publication bias was assessed using funnel plots.

Patient and public involvement
We did not seek patient or public comment in designing the study.

RESULTS
A total of 40 studies were included for analysis and their characteristics are summarised in online supplemental table S3. They comprised a total of 127 620 participants of whom 98 930 participants (77.5%) were classified as T1MI and 13 803 (10.8%) as T2MI. In the following text, we report key findings; more information and forest plots for each analysis involving more than one study and more than 100 total cases can be found in the online supplement, online supplemental figure S2–S4.

The 2007 definition was used in 7 (17.5%) studies, the 2012 definition in 25 (62.5%) studies, and the 2018 definition in 8 (20%) studies. Of the 40 studies, 17 (42.5%) were prospective and 23 (57.5%) were retrospective.
Of the 40 studies, 31 (77.5%) were assessed as good quality,21–26 27–33 34–39 40–46 as summarised in online supplemental table S4. Selection bias resulting in unrepresentative cohorts such as admission criteria to coronary care units or entry criteria into MI registries favouring T1MI,14 20 24–26 36 47 absence of independent adjudication of MI type as T1MI or T2MI,36 38 47 non-comparability of T1MI and T2MI cohorts,20 24 25 47 poorly specified outcome measures36 38 47 and short follow-up period resulting in few events14 20 24 36 comprised most forms of bias.

Funnel plots for in-hospital and 1-year all-cause mortality showed no asymmetry (online supplemental figures S45 and S46). Funnel plots for all other analyses showed similar results (available on request).

Participant characteristics

Patients with T1MI had a median age range of 60–82 years in the included studies that did not select a specific age population, compared with a median age range of 62–81 years in patients with T2MI. The sex distribution was also similar, with 58.4% and 53% of patients with T1MI and T2MI being male, respectively.

Regarding pre-existing medical conditions (table 1), patients with T2MI compared with patients with T1MI were more likely to have chronic kidney disease (22.8% vs 17.3%; OR 1.87; 95% CI 1.53 to 2.28), chronic heart failure (13.1% vs 7.6%; OR 2.35; 95% CI 1.82 to 3.03), atrial fibrillation (22.9% vs 6.1%; OR 3.02; 95% CI 2.29 to 3.99) and hypertension (66.4% vs 63.4%; OR 1.22; 95% CI 1.03 to 1.45). Patients with T2MI were less likely to have dyslipidaemia (43.4% vs 45.9%; OR 0.74; 95% CI 0.58 to 0.94) and smoking history (34.7% vs 52.8%; OR 0.6; 95% CI 0.49 to 0.73). There was no difference in the prevalence of type 2 diabetes mellitus or ischaemic heart disease between the two groups.

### Pre-existing medical conditions in patients with type 2 myocardial infarction (T2MI) versus type 1 myocardial infarction (T1MI)

| Pre-existing medical condition | T2MI | T1MI |
|-------------------------------|------|------|
| Number of patients with the specified condition | Total number of patients | % | Number of patients with the specified condition | Total number of patients | % | OR* (95% CI) |
| CAD | 3352 | 10 303 | 32.5 | 22 222 | 92 725 | 24 | 1.1 (0.93 to 1.31) |
| Type 2 DM | 3044 | 12 157 | 25 | 23 287 | 93 345 | 24.9 | 0.97 (0.85 to 1.10) |
| HTN | 7536 | 11 021 | 66.4 | 55 782 | 88 017 | 63.4 | 1.22 (1.03 to 1.45) |
| Dyslipidaemia | 4626 | 10 652 | 43.4 | 40 099 | 87 366 | 45.9 | 0.74 (0.58 to 0.94) |
| Smoker | 3448 | 9 929 | 34.7 | 39 548 | 74 889 | 52.8 | 0.60 (0.49 to 0.73) |
| Obesity | 1225 | 3 672 | 33.4 | 30 963 | 56 970 | 54.3 | 0.63 (0.46 to 0.87) |
| Chronic kidney disease | 1378 | 6 040 | 22.8 | 11 300 | 65 394 | 17.3 | 1.87 (1.53 to 2.28) |
| Heart failure | 1661 | 8 873 | 13.1 | 5 617 | 74 212 | 7.6 | 2.35 (1.82 to 3.03) |
| PVD | 584 | 5 856 | 10.0 | 2 066 | 41 280 | 5.0 | 1.33 (1.05 to 1.69) |
| CVD | 969 | 8 538 | 11.3 | 6 060 | 87 822 | 6.9 | 1.47 (1.27 to 1.71) |
| Atrial fibrillation | 836 | 3 645 | 22.9 | 1 220 | 19 843 | 6.1 | 3.02 (2.29 to 3.99) |
| COPD | 800 | 5 018 | 15.9 | 823 | 48 375 | 1.7 | 1.94 (1.22 to 3.08) |
| Illicit drug use | 46 | 204 | 22.5 | 8 | 220 | 3.6 | 8.15 (1.03 to 64.46) |

*Comparing patients with T2MI with those with T1MI, with OR adjusted according to study weighting using random effects meta-analysis.

**BMI, body mass index; CAD, coronary heart disease; COPD, chronic obstructive pulmonary disease; CVD, cerebrovascular disease; DM, diabetes mellitus; HTN, hypertension; PVD, peripheral vascular disease.**

**Risk-of-bias assessment**

Of the 40 studies, 31 (77.5%) were assessed as good quality,13 15–19 22 23 27–35 37–46 50–52 as summarised in online supplemental table S4. Selection bias resulting in unrepresentative cohorts such as admission criteria to coronary care units or entry criteria into MI registries favouring T1MI,14 20 24–26 36 47 absence of independent adjudication of MI type as T1MI or T2MI,36 38 47 non-comparability of T1MI and T2MI cohorts,20 24 25 47 poorly specified outcome measures36 38 47 and short follow-up period resulting in few events14 20 24 36 comprised most forms of bias. Funnel plots for in-hospital and 1-year all-cause mortality showed no asymmetry (online supplemental figures S45 and S46). Funnel plots for all other analyses showed similar results (available on request).

**Precipitating factors**

Less than half of the studies (n=18; 45%) included data on precipitating factors associated with T2MI.13 15 17 19 21–24 27 31 32 35 40 44 45 50–52 Data on each precipitating factor were not consistently available across the studies; for example, only 17 studies representing 45% of patients with T2MI assessed the presence of arrhythmia.

The most common precipitants were sepsis (35.9%) and heart failure (35.9%, followed by arrhythmia (29.8%) (online supplemental table S5), with non-cardiac surgery being deemed a cause in 12.2% of cases where data for this variable were collected.

**Presenting clinical features**

As summarised in online supplemental table S6, compared with patients with T1MI, patients with T2MI were less likely to present with typical cardiac symptoms of chest pain (38.6% vs 88.4%; OR 0.19; 95% CI 0.13 to 0.26) or discomfort in the arm or shoulder (8.5% vs 35%;
OR 0.18; 95% CI 0.11 to 0.3), but more likely to present with dyspnoea (27.1% vs 10.6%; OR 2.64; 95% CI 1.86 to 3.74).

**Investigations**

ECG findings on presentation (online supplemental table S7) such as ST elevation (14.1% vs 44.2%; OR 0.22; 95% CI 0.17 to 0.28) and pathological Q waves (6.7% vs 20.8%; OR 0.38; 95% CI 0.20 to 0.71) were less evident in T2MI than in T1MI. In contrast, non-specific ST-T wave changes (24.7% vs 10.8%; OR 2.64; 95% CI 1.86 to 3.74), and atrial arrhythmias (21% vs 6.6%; OR 4.99; 95% CI 3.14 to 7.93) were more common among T2MI. No differences between groups were seen in the frequency of ST depression or T wave inversion.

Among the 40 studies, 4 studies (10%) reported the use of high-sensitivity cTn assays, 21 (53%) reported sensitive assays and 14 (35%) did not specify what generation assay was used (online supplemental tables S3a and S3b). The results of troponin assays were reported in 26 (65%) studies, specific to cTnI assays in 19 studies, cTnT in 5, both assays in 1, while another did not specify the assay used. Only two of these studies reporting troponin failed to state the upper limit of normal (ULN) of the assay used.33 31 The troponin assays, and therefore units and reference ranges, varied between the studies, preventing direct comparison of troponin values. As a result, we converted troponin values to a multiple of the ULN for each assay to allow direct comparison (online supplemental table S8). For peak troponin, patients with T1MI had a higher and wider range of between 5 and 1702 times the ULN compared with patients with T2MI with a range of 2.8–447 times the ULN. Studies yielded mixed results as to whether the magnitude of change (or delta) in serial cTn assays was more predictive of T2MI or T1MI compared with absolute values of peak levels.33 Lowering the diagnostic threshold for troponin with the advent of more sensitive assays has increased the numbers of patients identified with T2MI by up to 50%,36 with more recent studies showing the incidence of T2MI equalling or exceeding that of T1MI.15 33 36

Echocardiography was less frequently performed among patients with T2MI than those with T1MI (47.9% vs 55.5%; OR 0.44; 95% CI 0.33 to 0.63) and when reported (online supplemental table S7), there was no difference in the prevalence of regional wall motion abnormalities or the level of left ventricular (LV) function, with reported median LV ejection fraction being 42.3%–55% in patients with T1MI and 40%–56% in patients with T2MI.

Coronary angiography was also less frequently performed among patients with T2MI than in those with T1MI (34.1% vs 85.5%; OR 0.09; 95% CI 0.06 to 0.12, online supplemental table S7). When performed, patients with T2MI were less likely to demonstrate obstructive coronary artery disease (34% vs 44.9%; OR 0.16; 95% CI 0.05 to 0.54), with obstruction variously defined as 50%–70% occlusion of one or more vessels.

**Management**

Patients with T2MI, compared with patients with T1MI, were significantly less likely to receive conventional cardioprotective medications (table 2), comprising beta-blockers (58.3% vs 76.3%; OR 0.45; 95% CI 0.33 to 0.63), antiplatelet agents (70.8% vs 88.5%; OR 0.25; 95% CI 0.16 to 0.38) and statins (52.9% vs 87.6%; OR 0.25; 95% CI 0.16 to 0.58). Of note, patients with T2MI were more likely to receive diuretics (44.8% vs 13.6%; OR 1.98; 95% CI 1.37 to 2.86).

**Table 2** Pharmacological management and invasive interventions in patients with type 2 myocardial infarction (T2MI) versus type 1 myocardial infarction (T1MI)

| Intervention | T2MI | T1MI |
|--------------|------|------|
| **Medication** | | |
| Beta blockers | 4967 | 8523 | 58.3 |
| ACEI/ARB | 3766 | 7842 | 48 |
| Antiplatelets | 5087 | 8599 | 70.8 |
| Anticoagulants | 1519 | 5255 | 28.9 |
| Antianginal agents | 1281 | 2191 | 58.5 |
| Diuretics | 1336 | 2985 | 44.8 |
| Statins | 3418 | 6455 | 52.9 |
| **Invasive** | | |
| PCI | 2092 | 9936 | 21.1 |
| CABG | 102 | 3451 | 2.9 |

Comparing patients with T2MI with those with T1MI, with OR adjusted according to study weighting using random effects meta-analysis. ACEI, ACE inhibitors; ARB, angiotensin receptor blockers; CABG, coronary artery bypass graft; PCI, percutaneous coronary intervention.

White K, et al. BMJ Open 2022;12:e055755. doi:10.1136/bmjopen-2021-055755

OR 0.18; 95% CI 0.11 to 0.3, but more likely to present with dyspnoea (27.1% vs 10.6%; OR 2.64; 95% CI 1.86 to 3.74).
White K, et al. BMJ Open 2022;12:e055755. doi:10.1136/bmjopen-2021-055755

Percutaneous coronary intervention (PCI) (21.1% vs 78.0%; OR 0.06; 95% CI 0.04 to 0.10) and coronary artery bypass surgery (2.9% vs 6.4%; OR 0.23; 95% CI 0.12 to 0.45) were also significantly less likely to be performed in patients with T2MI than patients with T1MI.

Prognosis

Patients with T2MI had significantly increased risk of all-cause death compared with patients with T1MI in both short-term and long-term follow-up (table 3). Specifically, compared with patients with T1MI, T2MI demonstrated increased all-cause mortality in-hospital (12.5% vs 5.8%; OR 1.94; 95% CI 1.35 to 2.79, online supplemental figure S40), at 1 year (18.9% vs 5.4%; OR 3.11; 95% CI 1.91 to 5.08, figure 1) and at 5–10 years (53.7% vs 28.5%, OR 3.24; 95% CI 2.73 to 3.84, figure 2). In contrast, there were no differences between patients with T2MI and T1MI in the risk of short-term mortality at 120–180 days (23.0% vs 12.5%; OR 1.34; 95% CI 0.63 to 2.85).

Table 3  Outcomes in patients with type 2 myocardial infarction (T2MI) versus type 1 myocardial infarction (T1MI)

| Outcomes                     | T2MI            | T1MI            | OR* (95% CI) |
|------------------------------|-----------------|-----------------|-------------|
| CV in-hospital mortality     | 184 2109 8.7    | 331 6248 5.3    | 1.61 (1.17 to 2.22) |
| All-cause in-hospital mortality | 667 5321 12.5  | 1508 25 997 5.8 | 1.94 (1.35 to 2.79) |
| Short-term all-cause mortality | 204 887 23.0  | 250 1998 12.5   | 1.34 (0.63 to 2.85) |
| 1-year all-cause mortality   | 632 3340 18.9   | 1299 24 203 5.4 | 3.11 (1.91 to 5.08) |
| 2-year all-cause mortality   | 246 926 26.6    | 428 2587 16.5   | 1.63 (1.11 to 2.41) |
| 3-year all-cause mortality   | 193 525 36.8    | 710 4305 16.5   | 2.00 (1.07 to 3.76) |
| Long-term all-cause mortality| 1453 2708 53.7  | 1320 4633 28.5  | 3.24 (2.73 to 3.84) |

*Comparing patients with T1MI with those with T2MI, with OR adjusted according to study weighting using random effects meta-analysis. CV, cardiovascular.

DISCUSSION

To our knowledge, this is the most comprehensive systematic review and meta-analysis of contemporary studies comparing T2MI with T1MI in the troponin era, comprising 127 620 patients from 40 cohort studies across 14 countries, and which used formal definitions of T2MI and T1MI. Up to three quarters of all MIs in routine care can be T2MI, and distinguishing T2MI from T1MI on clinical criteria is often challenging. The management strategies used by clinicians in real-world practice for T2MI often vary, and the clinical outcomes of T2MI compared with T1MI, particularly over the long term, have been uncertain. This review provides information that helps characterise these two groups of patients according to multiple variables and which may assist in clinical decision-making and prognostication.

In this review, patients with T2MI demonstrated more medical comorbidities than patients with T1MI, as noted in a recent meta-analysis. Our review highlighted the much higher incidence of pre-existing generalised vascular disease, atrial fibrillation, renal impairment and heart failure among patients with T2MI.

Figure 1  Forest plot of 1-year all-cause mortality of patients with type 2 myocardial infarction (T2MI) compared with patients with type 1 myocardial infarction (T1MI).
Figure 2 Forest plot of long-term all-cause mortality of patients with type 2 myocardial infarction (T2MI) compared with patients with type 1 myocardial infarction (T1MI).

Sepsis\textsuperscript{10, 16, 27} and anaemia\textsuperscript{51} ranked highly as triggers, together with other acute cardiac events such as valve dysfunction or arrhythmias. In one study, a more favourable prognosis in T2MI was seen when the principal trigger was arrhythmia compared with non-cardiac surgery, hypotension, anaemia or hypoxia.\textsuperscript{29} In another study, shock syndromes were triggers portending a worse prognosis compared with all other triggers.\textsuperscript{32} In our analysis, non-cardiac surgery as a trigger was less frequent than reported by other investigators\textsuperscript{26} whereby perioperative stressors including blood loss, anaesthesia-induced hypotension and wound infections cause imbalance in myocardial contractility, oxygen demand and blood flow.\textsuperscript{53}

Analysis of cTn levels showed uniformly higher values in T1MI than T2MI which accord with one review\textsuperscript{5} reporting cTn values 30\%–94\% higher in patients with T1MI, and which other investigators regard as being highly specific diagnostic markers for T1MI.\textsuperscript{53}

Coronary angiography and revascularisation were both performed much less frequently in patients with T2MI than those with T1MI. Treating physicians may perceive invasive strategies as being contraindicated or potentially harmful in the presence of various comorbidities more commonly seen in T2MI and associated with competing mortality risk. In our pooled data, only one in three patients with T2MI who underwent angiography demonstrated obstructive coronary artery disease, although this figure may be an underestimate due to selection bias whereby younger, less multimorbid patients preferentially underwent angiography. In the Catheter Sampled Blood Archive in Cardiovascular Disease (CASABLANCA) cohort study, which enrolled patients with high likelihood of coronary or peripheral artery disease and subjected them to peripheral or coronary angiography, of all those who subsequently suffered incident T2MI, almost half (47.7\%) demonstrated \( \geq 70\% \) stenosis in at least 2 major coronary arteries.\textsuperscript{54} These conflicting findings question whether patients presenting with T2MI would benefit from routine use of invasive strategies that define coronary anatomy and, if plaque rupture or critical stenoses are seen, lead to prompt revascularisation, with resultant improvement in patient outcomes. In one study,\textsuperscript{18} "angiography unmasked acute plaque rupture in 29\% of patients classified as T2MI. In another study, among 27 of 236 patients with T2MI who underwent revascularisation, the odds of all-cause death were reduced by 67\% compared with the remaining 209 non-revascularised patients.\textsuperscript{23} In contrast, in a third more rigorous study comparing patients with T2MI versus patients with T1MI who received or did not receive PCI within 24 hours of symptom onset, after adjusting results using multivariate logistic regression analysis and inverted probability weighting,\textsuperscript{15} in-hospital mortality was lower in those with T1MI receiving PCI (OR 0.47; 95\% CI 0.40 to 0.55; \( p<0.001 \)), but not in those with T2MI receiving PCI (OR 1.09; 95\% CI 0.62 to 1.94; \( p=0.763 \)). However, all these studies are observational, so completion of randomised trials, such as the appropriate-ness of coronary investigation in myocardial injury and T2MI (ACT-2) trial, which is currently in recruitment,\textsuperscript{55} will hopefully provide a more definitive answer.

Given that a third of patients with T2MI had pre-existing coronary artery disease and most of the remainder had one or more cardiovascular risk factors, the relative underuse of cardioprotective medications is perplexing. It may reflect either clinician uncertainty around their cardioprotective utility in T2MI, or concerns about the potential for adverse interactions with other drugs or diseases commonly seen in multimorbid patients with T2MI. The higher use of diuretics in the T2MI population likely reflects the higher prevalence of heart failure and hypertension. Recognising the heterogeneous mechanisms or conditions leading to T2MI, a phenotype specific approach to the design of future trials will be useful in identifying effective therapies.

An important finding is the much higher all-cause in-hospital and 1-year mortality in patients with T2MI compared with patients with T1MI, similar to the twofold greater mortality rate in T2MI noted in a recent systematic review of nine studies.\textsuperscript{8} In our review, this excess mortality was not driven by an excess of cardiovascular deaths, and likely reflects the competing risks of multiple comorbidities, rather than underlying obstructive coronary artery disease which was seen in 30\%–50\% of patients with T2MI.\textsuperscript{26} Studies yielded mixed results as to whether coronary artery disease is an independent predictor of T2MI,\textsuperscript{26, 42} while others question the angiographic distinction between T2MI and T1MI. For example, in a study of 450 consecutive patients with MI who all underwent coronary angiography within 24 hours of symptom onset, 145 (32.2\%) patients had ‘true’ T1MI
(acute atherothrombosis and no systemic triggers), 114 (25.3%) had ‘true’ T2MI (no atherothrombosis and systemic triggers), 61 (13.6%) patients had neither, and 130 (28.9%) patients had both. This yielded a discordance of angiographic and clinical definitions of MI type in 42.5% of patients.

Our review has several limitations. First, in the absence of individual patient data from all included studies, we could not perform multivariate regression analysis in identifying independent predictors of diagnosis, management, or prognosis of T2MI. Second, we did not perform separate analyses of studies according to each version of the universal definition of MI or to different troponin thresholds to define MI, which may impact management and prognosis. However, potential misclassification bias was addressed in a recent study which showed little change in MI classification as type 1 or 2 in the same cohort of emergency admissions to whom the third and fourth universal definitions were applied. In another study which compared separate T2MI cohorts, as defined by the 2007 and the 2012 definitions, comorbidities and use of cardioprotective medications were less frequent in the 2012 cohort, likely due to less severe MI being included as a result of the use of more sensitive troponin assays. Third, we did not collect haemodynamic variables or other physiological measures such as haemoglobin levels and glomerular filtration rate in analysing clinical presentations as these were very inconsistently reported. Fourth, our mortality meta-analyses relied on crude mortality rates reported in each study, with 57% of studies also undertaking multivariate regression and/or competing risk analyses and reporting adjusted mortality rates. For the T2MI cohorts in general, these rates tended to be lower and the differences in rates compared with those of T1MI were of smaller magnitude. Similarly, we did not attempt subanalyses based on risk stratification using validated risk scores or seek to identify predictive models for mortality, as such analyses were reported in only two studies. Fifth, we did not analyse 30-day readmission rates as these were reported in only three studies. Sixth, we did not perform sensitivity analyses comparing results of prospective vs retrospective studies, as neither group demonstrated less or more risk of bias than the other, or compared results of good quality studies against fair/poor quality studies as the latter comprised only 17% of all patients. Seventh, as we searched only two databases and did not include grey literature, relevant studies may have been missed. However, in a recent analysis, searching Medline and Embase combined yielded 93% of relevant studies, with Google Scholar, despite requiring much more time and effort, only yielding another 3%. Eighth, while publication bias is possible, all funnel plots performed for every analysis showed no asymmetry. Finally, we did not perform subgroup analyses or meta-regression in assessing between-study heterogeneity, as study parameters (such as study design and analytic methods) were often ill-defined and widely variable across this large number of real-world observational studies.

The strengths of this review are the inclusion of all contemporary cohort studies in the troponin era that employed formal definitions of T2MI, analysis of a broader range of variables than those of previous studies, and the more precise discernment of clinically meaningful differences between the two MI populations in patient characteristics, clinical presentation, patterns of care and outcomes. As studies originated from different jurisdictions, we believe our findings are generalisable to different healthcare systems, although absolute values for some measures did vary between countries. We are aware of a large US cohort study published since completion of our review which compared patients with T1MI with those with T2MI, but was limited by misclassification bias (relying on administrative hospital discharge data containing an International Classification of Diseases-10th Revision code specific for type 2 MI, rather than a registry or chart diagnosis based on a formal MI definition), short study period of 3 months in late 2017, and inability to analyse clinical features, investigation results, medication use, coronary anatomy, and postdischarge mortality due to their omission in the datasets.

CONCLUSION

This review has identified differences between patients with T2MI and T1MI in presenting clinical features, investigation and management profiles and clinical outcomes. These findings may assist clinicians to better recognise T2MI and advise patients about its sequelae, and inform hospital coding and epidemiological trending, quality of care indicators, and interhospital benchmarking of performance related to the care of patients with T2MI.

The review has also defined persisting gaps in our understanding of the utility and prognostic effects of invasive investigations, revascularisation strategies and cardioprotective medications in patients with T2MI that warrant more randomised trials that enrol such patients.

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Data availability statement All data relevant to the study, except for funnel plots for non-mortality analyses, are included in the article or uploaded as supplementary information. Funnel plots not included are available upon request.
Supplemental material

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