Increased cancer risk after myocardial infarction: fact or fiction? A systemic review and meta-analysis

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Objective: Accumulating evidences showed some positive relations between myocardial infarction (MI) and new onset cancer. We aim to investigate whether MI is associated with an increased risk of incident cancer.

Methods: A comprehensive literature list was identified from MEDLINE, Embase, and Web of Science databases from inception until October 2018. The main inclusion criteria included observational studies investigating the association between MI and new onset cancer. Stata 12.0 software was used for meta-analysis.

Results: Of 862 potentially relevant studies, five cohort studies met all inclusion criteria. The pooled cancer incidence rate was 9.5% (95% CI=8.3–10.7%). Pooled analysis of OR showed that the increased overall cancer risk in MI patients in comparison with controls had no statistically significant (OR=1.08; 95% CI=0.97–1.19, P=0.153). Subgroup analysis by gender demonstrated that the overall cancer risk was only significantly increased in female (OR=1.10; 95% CI=1.01–1.20, P=0.025), but not in male patients (OR=1.04; 95% CI=0.99–1.10, P=0.124). In terms of cancer type, the increased cancer risk was only significant for lung cancer (male OR=1.12; 95% CI=1.05–1.19, P<0.01; and female OR=1.51; 95% CI=1.15–1.99, P<0.01), but not for prostate (OR=0.96; 95% CI=0.85–1.09, P=0.546) or breast cancer (OR=0.94; 95% CI=0.86–1.04, P=0.222). In addition, the increased cancer risk was only significant in the first 6 months (OR=1.93; 95% CI=1.42–2.63, P<0.01) but not in 6 months–1 year (OR=1.03; 95% CI=0.92–1.15, P=0.627) or >1 year (OR=0.98; 95% CI=0.93–1.04, P=0.585) follow-up after MI.

Conclusion: From available evidence, the increased overall cancer risk after MI was only significant in female but not in male patients. Besides, the increased cancer risk could be driven by increased short-term cancer incidence after MI and certain cancer types such as lung cancer.

Keywords: myocardial infarction, incident cancer, meta-analysis

Introduction
Myocardial infarction (MI) and cancer are two leading causes of morbidity and mortality worldwide. With the development of pharmacotherapy and timely cardiac revascularization, in past decades, the prognosis of MI has been fundamentally improved with higher survival rate.1 Recently, some studies have suggested cardiovascular diseases including MI and cancer share a number of risk factors and possess various similarities.2,3 Inflammation, oxidative stress, obesity, and smoking are all contributors to the occurrence of MI and cancer.4,7 Some observational studies also found noncardiac causes (mainly malignancies and chronic diseases) are responsible for the majority of later deaths in MI patients treated with percutaneous coronary intervention.8,9 Thus, it...
is plausible to assume that patients with MI history are more likely to develop incident cancer.

Meanwhile, in past years, several clinical and epidemiological studies have been done in this field and showed some positive associations between MI and new onset cancer.\(^{10,11}\) However, the results were inconsistent or even conflicting.\(^{12,13}\) In view of the high disease burden of MI and cancer, it is necessary to confirm whether MI is a new risk factor for incident cancer. Therefore, we conducted a systematic review and meta-analysis in accordance with the PRISMA guidelines and our predesigned protocol to clarify the association between MI and incident cancer.

## Methods

### Literature search

Two investigators independently reviewed published studies in MEDLINE, Embase, and Web of Science databases from their inception to October 2018 by using the following index terms: myocardial infarction OR myocardial infarct OR heart attack OR myocardial injury AND cancer incidence OR incident cancer OR new onset cancer OR cancer risk OR carcinogenesis OR tumorigenesis OR oncogenesis. A manual search for additional studies using references of selected retrieved articles was also performed to identify other possible studies. We also conducted searches of conference proceedings from major cardiology and oncology meetings for additional abstracts on the topic. No limitation on language was applied. The last date of the search was October 23, 2018.

### Inclusion and exclusion criteria

The inclusion criteria were as follows: 1) observational studies; 2) studies investigating myocardial infarction and cancer incidence; 3) myocardial infarction was clearly diagnosed and new onset cancer occurred after myocardial infarction; and 4) cancer incidence rate or RR, OR, HR or standardized incidence ratio (SIR) with 95% CI were provided.

Exclusion criteria: 1) duplicate publications; 2) publication types were not observational studies; 3) cancer incidence was before or concurrent with MI; and 4) studies were also excluded if relevant data were not reported or extractable.

### Quality assessment

The quality of each study was assessed by two reviewers. Newcastle Ottawa scale (NOS) generating stars for selection (S), comparability (C) and outcome (O) for cohort and case-control studies were used to evaluate studies’ quality and risk of bias.\(^{14}\) Studies with seven stars or more were assumed to be of high quality, between five and seven stars to be of moderate quality, and less than five stars to be of low quality.

### Data extraction

Data extraction was performed independently using a predefined data extract form by two reviewers. The data extracted from each study included the following information: first author’s name, publication year, study design, country, number of patients and controls, gender, cancer incidence, and effect estimates (OR, RR, HR, or SIR) with 95% CI and factors adjusted in the multivariate analysis. If both crude and adjusted values were provided, we only extracted the adjusted ones. Disagreement was resolved by discussion among all investigators.

### Statistical analysis

All analysis was performed using Stata 12.0 software (StataCorp, College Station, TX). The risk of cancer incidence after MI was assessed by ORs and the corresponding 95% CIs. An OR <1 reflects a favorable outcome in MI group compared with controls and indicates lower cancer incidence rate, and vice versa. A two-sided \(P\)-value of 0.05 was considered statistically significant for all included analysis. Generic inverse variance (DerSimonian and Laird) method was employed to combine adjusted point estimates and standard errors from each study.\(^{15}\) If effect estimates were presented for a number of categories of exposure, we would combine the corresponding estimates using the method proposed by Hamling et al.\(^{16}\)

\(Q\) test and \(I^2\) statistic were used to ascertain the between-study heterogeneity. A value of \(I^2\) of 0%–25%, 25%–50%, 50%–75%, and >75% embodied insignificant, low, moderate, and high heterogeneity, respectively.\(^{17}\) In light of the high likelihood of between-study variance, we used a random-effect model rather than a fixed-effect model. Subgroup analysis would be done based on gender, cancer type, and follow-up time.

Publication bias was assessed by Begg’s test and Egger’s test.\(^{18,19}\) \(P<0.05\) indicated bias, and \(P>0.05\) indicated no publication bias.

### Results

#### Search results and characteristics of studies included

Of 862 potentially relevant articles, 846 articles were excluded due to duplication or the title and abstract not meeting inclusion criteria. Subsequently, 11 articles were excluded because of data duplication or lack of the outcomes
of interest. Finally, five cohort studies met all inclusion criteria.10–13,20 One of them was a single-arm cohort study regarding MI and only provided cancer incidence rate. The rest of the studies supplied overall or subgroup effect estimates (OR, RR, HR, or SIR) with 95% CI. The literature retrieval, review, and selection process are shown in Figure 1. The characteristics of cohort studies are shown in Table 1. The quality assessment of observational studies was performed by NOS quality scale (Table 1). Quality assessment results suggested observational studies included had moderate or high quality, which might lead to heterogeneity and risk of bias.

**Meta-analysis results**

### Cancer risk after MI

Four studies with 221,994 MI patients were included to estimate the incidence rate of cancer among patients with MI history.10,12,13,20 The pooled cancer incidence rate in MI patients was 9.5% (8.3%–10.7%, Figure 2), which was comparable to that in non-MI controls (11.1%; 95% CI=7.3%–14.8). Three studies reported the overall effect estimates in patients with MI.10,12,13 Pooled analyses of OR in the random-effects model showed that the increased cancer risk in patients with MI in comparison with controls had no statistical significance, and the heterogeneity among studies was high (OR=1.08; 95% CI=0.97–1.19, P=0.153; I²=90.8%; P for heterogeneity <0.01; Figure 3).

#### Subgroup analysis by gender

Subgroup effect estimates based on gender were presented in four studies.10–13 Pooled analyses of OR in random-effects model demonstrated that the overall risk of cancer incidence in male patients with MI was nonsignificant (OR=1.04; 95% CI=0.99–1.10, P=0.124; I²=78.5%; P for heterogeneity <0.01; Figure 4A). However, female patients with MI had a significantly increased risk of incident cancer (OR=1.10; 95% CI=1.01–1.20, P=0.025; I²=82.9%; P for heterogeneity <0.01; Figure 4B).

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**Figure 1** Flow diagram of literature review process.
Table 1 Characteristics of studies included

<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Country</th>
<th>Design</th>
<th>No. of MI patients</th>
<th>Age (years)</th>
<th>Male proportion (%)</th>
<th>OR or equivalents with 95% CI</th>
<th>Follow-up time (years)</th>
<th>Adjusted factors</th>
<th>Quality assessment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hasin et al</td>
<td>2016</td>
<td>USA</td>
<td>Cohort</td>
<td>1,081</td>
<td>64±15</td>
<td>60</td>
<td>NA</td>
<td>4.9±3.0</td>
<td>NA</td>
<td>Moderate</td>
</tr>
<tr>
<td>Pehrsson et al</td>
<td>2005</td>
<td>Sweden</td>
<td>Cohort</td>
<td>NA</td>
<td>&lt;80</td>
<td>65.20</td>
<td>Men: 1.08 (1.04–1.11); Women: 1.15 (1.09–1.21)</td>
<td>9.3 (0–28)</td>
<td>Age</td>
<td>High</td>
</tr>
<tr>
<td>Dreyer and Olsen</td>
<td>1998</td>
<td>Denmark</td>
<td>Cohort</td>
<td>96,891</td>
<td>Men: 63; Women: 69</td>
<td>67.97</td>
<td>1.05 (1.03–1.07)</td>
<td>5.9 (1–17)</td>
<td>NA</td>
<td>Moderate</td>
</tr>
<tr>
<td>Rinde et al</td>
<td>2017</td>
<td>Norway</td>
<td>Cohort</td>
<td>1,747</td>
<td>62±13</td>
<td>62.0</td>
<td>1.46 (1.21–1.77)</td>
<td>Median: 15.7</td>
<td>Systolic blood pressure, diabetes mellitus, HDL, smoking, physical activity, and education level</td>
<td>High</td>
</tr>
<tr>
<td>Malmborg et al</td>
<td>2018</td>
<td>Denmark</td>
<td>Cohort</td>
<td>122,275</td>
<td>Men: 59.2 (49.5–69.5); Women: 68.5 (58.1–76.0)</td>
<td>61.2</td>
<td>0.97 (0.92–1.01)</td>
<td>0–17</td>
<td>Age, sex, calendar year, hypertension, dyslipidemia, diabetes, COPD and socioeconomic status</td>
<td>High</td>
</tr>
</tbody>
</table>

Abbreviations: HDL, high-density lipoprotein; MI, myocardial infarction.

Study ID | ES (95% CI) | Weight |
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Hasin 2016</td>
<td>0.09 (0.07, 0.11)</td>
<td>18.27</td>
</tr>
<tr>
<td>Dreyer 1998</td>
<td>0.11 (0.11, 0.11)</td>
<td>29.86</td>
</tr>
<tr>
<td>Rinde 2017</td>
<td>0.08 (0.07, 0.10)</td>
<td>21.94</td>
</tr>
<tr>
<td>Malmborg 2018</td>
<td>0.09 (0.09, 0.09)</td>
<td>29.94</td>
</tr>
<tr>
<td>Overall (P=98.0%, P=0.000)</td>
<td>0.10 (0.08, 0.11)</td>
<td>100.00</td>
</tr>
</tbody>
</table>

Note: weights are from random effects analysis

Figure 2 Forest plot of cancer incidence rate after myocardial infarction.
Subgroup analysis by cancer type
Three studies reported the outcome of cancer types or sites. Although there was much difference in the classification of cancers in these studies, all of them provided the gender-stratified incidence of lung, prostate, and breast cancers. We found lung cancer risk was significantly increased both in male (OR = 1.12; 95% CI = 1.05–1.19, P < 0.01; I² = 38.6%; P for heterogeneity = 0.196; Figure 5A) and female (OR = 1.51; 95% CI = 1.15–1.99, P < 0.01; I² = 92.0%; P for heterogeneity < 0.01; Figure 5B) patients with MI history. However, the increased risk of sex-specific cancers including prostate cancer (OR = 0.96; 95% CI = 0.85–1.09, P = 0.546; I² = 94.4%; P for heterogeneity < 0.01; Figure 5C) and breast cancer (OR = 0.94; 95% CI = 0.86–1.04, P = 0.222; I² = 39.9%; P for heterogeneity = 0.189; Figure 5D) had no statistical significance.

Subgroup analysis by follow-up time
Only two studies revealed the incidence of cancer in subjects with MI by time after the MI event. We observed the increased cancer risk was only significant in the first 6 months after MI with no heterogeneity (OR = 1.93; 95% CI = 1.42–2.63, P < 0.01; I² = 0%; P for heterogeneity = 0.606; Figure 6A). After 6 months, the increased cancer risk vanished, which was reflected by the pooled ORs for 6 months to 1-year (OR = 1.03, 95% CI = 0.92–1.15, P = 0.627; I² = 0%; P for heterogeneity = 0.786; Figure 6B) and >1-year (OR = 0.98, 95% CI = 0.93–1.04, P = 0.585; I² = 0%; P for heterogeneity = 0.487; Figure 6C) follow-up after MI.

Evaluation for publication bias
Begg’s and Egger’s tests were performed to evaluate publication bias. For the overall cancer risk analysis of MI vs control, there was no significant publication bias (Begg’s test, P = 0.602; Egger’s test, P = 0.816).

Discussion
To our knowledge, this is the first systematic review and meta-analysis of published observational studies assessing the association between MI and incident cancer. Our results showed that the estimated cancer incidence rate after MI was 9.5% and the relation between MI and cancer incidence was uncertain. The pooled ORs of increased overall cancer risk were only significant in female patients, but not in male patients. Although the OR of cancer risk in female patients reached statistical significance, the absolute increased risk was only 10% (OR = 1.10; 95% CI = 1.01–1.20, P = 0.025) and the heterogeneity was high. In terms of cancer type, the increased cancer risk was only significant for lung cancer, but not for prostate cancer or breast cancer. In addition, subgroup analysis by follow-up time suggested the increased risk of incident cancer did not persist over 6 months after MI event.

Sharing risk factors are one of the important reasons for association between MI and cancer. These shared risk factors can contribute to both the cardiovascular and the malignant event, with the malignancy occurring later. As a result, chance is a possible explanation for the increased cancer incidence after MI, and the independent association...
between MI and cancer may have been overlooked. Besides, the shared risk factor ‘smoking’, a known lung cancer cause, may partially account for our observation that lung cancer risk was significantly increased after MI.

In addition, the time since MI diagnosis must be considered when investigating cancer incidence. It is necessary to do such estimates by time, as cancer usually develops and evolves over several years. If cancer incidence is observed shortly after the start of MI follow-up, MI would be less likely to be a causal factor, and occult cancers could have occurred before the cardiovascular event. In our analysis, we did a subgroup analysis by follow-up time and found an interesting phenomenon. The risk of cancer incidence was the highest in the first follow-up period (<6 months). With follow-up time increasing, the ORs decreased sharply and became nonsignificant. This suggested that the higher cancer risk

**Figure 4** Forest plots of overall cancer risk after myocardial infarction by gender.

Note: (A) male patients; (B) female patients.
**Figure 5** Forest plots of cancer risk after myocardial infarction by cancer types.

**Note:** (A) lung cancer in male patients; (B) lung cancer in female patients; (C) prostate cancer in male patients; (D) breast cancer in female patients.
Figure 6 Forest plots of overall cancer risk after myocardial infarction by follow-up time.
Note: (A) <6 months; (B) 6 months–1 year; (C) >1 year.
risk shortly after MI may not be due to MI itself, but other confounding factors such as surveillance bias. Patients with MI had more frequent clinical encounters with more diagnostic tests, especially in the first several months after MI event, and this would increase the chance of early detection of cancer.13

Although the above-mentioned shared risk factors, occult cancers, and surveillance bias may explain the increased cancer incidence after MI, we couldn’t exclude the possibility that MI itself can cause a higher risk of long-term cancer development. A recent large cohort study presented that, after a median follow-up of 1,020 days, atherosclerotic cardiovascular disease (ASCVD) itself increased the risk of cancer incidence.23 On the other hand, a laboratory experimental study by Meijers et al24 just showed heart failure stimulates tumor growth by cardiac excreted circulating factors. MI belongs to ASCVD and can also lead to heart failure in some patients. Therefore, it is still possible that MI can have some effect on the long-term risk of cancer incidence. As Hasin et al21 summarized, malignancy may be caused by biological alterations or treatment modalities related to the cardiovascular diseases. In the future, the links between MI and new onset cancer remain to be established by more basic and clinical research.

Limitations
There are several limitations in our study. First, the present number of studies on MI and incident cancer is limited, especially in the investigation of cancers types (three studies) and cancer incidence by time (two studies). Since only two available studies were included in the analysis of cancer risk by follow-up time, the results should be interpreted with caution. Second, our study is related to the observational nature of the studies included with all inherent biases of observational designs. Third, heart failure also has some links with cancer incidence, but most of the studies included did not supply data on heart failure or left ventricular ejection fraction. Therefore, heart failure could become a confounder. However, current evidence represented the best available and all studies included were of moderate-to-high quality, including population-based studies.

Conclusion
From available evidence, the increased overall cancer risk after MI was only significant in female but not in male patients. Besides, the increased cancer risk could be driven by increased short-term cancer incidence after MI, and certain cancer types, such as lung cancer. However, due to the limitations listed above, further studies with larger sample sizes and long-term follow-up are warranted to establish the association between MI and new onset cancer.

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Disclosure
The authors report no conflicts of interest in this work.

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