Primary Headache and Silent Myocardial Ischemia in Patients with Coronary Artery Disease

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Abstract

Objective: The mechanisms by which migraine is linked to ischemic vascular disease remain uncertain and are likely to be complex. The aim of this study was to investigate the correlation between silent myocardial ischemia (SMI) and a history of documented primary headache in a large population of patients with exercise-induced myocardial ischemia.

Methods: The study involved 1,427 consecutive patients (918 symptomatic and 509 asymptomatic patients) with exercise-induced myocardial ischemia and documented coronary artery disease (CAD).

Results: Patients with anginal symptoms during exercise-induced myocardial ischemia had a significantly higher prevalence of primary headache than those without (41 vs. 30%, p < 0.001). Patients with angina pectoris in daily life also had greater prevalence of primary headache than those without anginal symptoms (37 vs. 20%; p < 0.0001). Symptomatic patients during percutaneous transluminal coronary angiography or myocardial infarction had a greater prevalence of primary headache than asymptomatic patients (p < 0.001 and p = 0.005, respectively). Conclusions: Our data suggest that a history of headache in CAD population is correlated to a high probability of anginal symptoms and a decreased probability of SMI. The anamnestic absence of headache requires a close monitoring for patients with risk factors for CAD, because this population seems to have a lower susceptibility to pain and the risk of developing SMI might be increased.

Introduction

Silent myocardial ischemia (SMI) is defined as an objective documentation of myocardial ischemia in the absence of angina or anginal equivalents [1].

According to the Cohn classification [2], SMI can occur in: (i) totally asymptomatic individuals without any...
evidence of known coronary artery disease (CAD), (2) subjects who are asymptomatic after having a myocardial infarction (MI) and (3) those with both symptomatic and asymptomatic ischemic episodes. Groups 2 and 3 can be merged so that group 1 consists of patients without known CAD and group 2 with known disease [3]. This latter group is at particularly high risk for cardiovascular events [4–6].

The presence or absence of anginal symptoms may be partly explained by individual differences in the pain threshold [7–9]. In addition, chemical factors released in response to ischemia which stimulate cardiac afferent pain fibers may be responsible for cardiac pain; various chemical agents have been implicated as a stimulus for angina pectoris during ischemia [10, 11]. However, the mechanism underlying SMI is not well understood, and many theories have been advanced.

Headaches are most thoroughly classified by the International Headache Society’s International Classification of Headache Disorders [12, 13], which divides headaches into two broad categories: primary headache disorders (which include migraine, tension-type and cluster headache) and secondary headache.

Some studies have explored the possible association between primary headache and CAD, especially in subjects with migraine [14–17]. In addition, migraine, especially with aura, is an established risk factor for ischemic lesions of the brain [18]. Recent evidence has linked migraine to a broader range of ischemic vascular disorders, including angina, MI, coronary revascularization, claudication and cardiovascular mortality [19]. The mechanisms by which migraine is linked to ischemic vascular disease remain uncertain and are likely to be complex. To the best of our knowledge, there are no data available in the literature regarding the association between primary headache and SMI. Therefore, the aim of this study was to explore whether the absence of anginal symptoms during myocardial ischemia was associated to reduced sensibility to pain. In particular, we investigate the possible correlation between SMI and a history of primary headache (migraine, tension-type and cluster headache) and secondary headache.

Methods

Patient Selection

This study included 1,427 consecutive patients (1,188 men and 239 women) who underwent both an exercise test and coronary angiography and presented the following characteristics: (1) exercise-induced myocardial ischemia (ischemic electrocardiographic response of horizontal or downsloping ST-segment depression of at least 0.1 mV during exercise testing) and (2) angiographically documented significant CAD defined as the presence of greater than 50% narrowing of the luminal diameter of one or more coronary arteries.

Patients with pain syndrome or acute pain at the time of examination were excluded from the study as well as patients with cerebrovascular events, trauma or known cognitive, sensory and motor defects.

The study protocol conformed to the ethical guidelines of the Declaration of Helsinki. Informed written consent was obtained from all subjects and the study was approved by the ethical committee. The authors of this manuscript have certified that they comply with the principles of ethical publishing in the International Journal of Cardiology [20].

Exercise Stress Test

Exercise stress testing was performed as previously described [6]. Briefly, a multistage-bicycle-ergometer exercise stress test was performed with the patients in the supine position with an initial workload of 25 W and successive increments of 25 W every 3 min at a pedaling frequency of 60 rpm. A standard 12-lead electrocardiography was taken before, at the end of each stage and at the end of exercise. Leads V4, V5, V6 were monitored during exercise. The systolic and diastolic blood pressures were measured with a sphygmomanometer at basal conditions, at the end of each interval, at the maximum amount of work and every 3 min in the recovery phase. During the test, the patients were interrogated continuously regarding their symptoms in terms of pain, dyspnea, vertigo and tachycardia among others. Patients who always reported anginal symptoms were enrolled in the symptomatic group and those who did not complain of anginal pain were enrolled in the asymptomatic group. Patients with pacemaker, valvular disease or heart failure were excluded from the study.

The test was interrupted if one of these conditions manifested: fatigue, appearance of chest pain and/or dyspnea, exhaustion, ST-segment elevation ≥1 mm without diagnostic Q waves, ST-segment depression ≥2 mm, repetitive arrhythmias and a drop in systolic blood pressure >10 mm Hg. The test was considered positive in the presence of downsloping ST-segment depression ≥1 mV below the baseline, at least 0.06–0.08 s after the QRS complex J-point in multiple leads or ST-segment elevation.

Before the exercise test, pharmacological washout was performed. Calcium-channel antagonists and nitrates were halted 48 h before the test. β-Blockers were gradually reduced and then stopped 1 week before the examination. Patients who were taking digitalis were not included in this study.

Cardiovascular Risk Factors

Diabetes was diagnosed according to the American Diabetes Association criteria [21]. Hypertension was diagnosed according to the European Society of Hypertension/European Society of Cardiology criteria [22] or in the presence of a specific treatment. A family history of CAD was defined as documented evidence of premature CAD in a close relative (men <55 and women <65 years of age). Subjects were classified as currently smoking, as having stopped smoking in the past (if they had stopped >4 weeks and <40 years earlier), or as never having smoked (if they had never smoked or had stopped ≥40 years earlier). All patients completed the Perceived Stress Scale [23].
Coronary Angiography

All subjects underwent diagnostic coronary angiography using standard techniques, as previously described [6]. CAD was defined as angiographic evidence of stenosis in any epicardial coronary artery of ≥50% of its diameter. CAD extent was defined by the number of major epicardial coronary vessels with angiographic evidence of one or more stenosis of ≥50% severity.

Ascertainment of Headache Status

Trained interviewers asked participants about their lifetime history of headaches based on the criteria of the second edition of the International Headache Society’s Classification of Headache Disorders which represents the most important document for the diagnosis and management of headache patients. This classification divides headaches into 3 categories: (1) primary, (2) secondary and (3) cranial neuralgias, facial pain and other headaches. In particular, we looked at the presence of tension-type, migraine or cluster headaches, which are the three most common forms of primary headache [13].

Statistical Analysis

The Kolmogorov-Smirnov test of normality was used to verify whether the distribution of variables followed a Gaussian pattern. Normally distributed data in groups were expressed as means ± SDs. For continuous variables, the differences between the groups were evaluated with an unpaired Student t test. To explore the independence between SMI and several variables, a multiple regression analysis was performed. Nonnormally distributed variables were log-transformed before the analyses. Categorical variables are presented by frequency counts, and intergroup comparisons were analyzed by a χ² test. Two-tailed p < 0.05 was considered statistically significant.

Results

Table 1 shows clinical, biological and angiographic feature of the whole population with exercise-induced myocardial ischemia and patients stratified by the presence/absence of anginal symptoms. Patients with symptoms during exercise-induced myocardial ischemia had a significantly greater prevalence of primary headache than those without (41 vs. 30%; p < 0.001).

Multivariate Analysis

To assess whether the association between primary headache and SMI was independent, a multiple regression analysis was performed with the following variables as potential predictors: hypertension, dyslipidemia, a positive family history of CAD, ever-smoking, stress, diabetes, BMI, age and medical treatment. In the multiple regression analysis, we found that primary headaches were independently associated with angina (β = 0.095; p < 0.001).

Subgroup Analysis

Diabetic patients who had chest pain during the exercise-stress test had a greater prevalence of primary headache than asymptomatic diabetic patients (45 vs. 27%; p = 0.002). Likewise, in symptomatic hypertensive patients, the prevalence of primary headache was greater than in asymptomatic hypertensive patients (44 vs. 27%; p < 0.0001).

We also analyzed the correlation between a history of primary headache and episodes of anginal symptoms during daily life, PTCA and MI as shown in figure 1. The presence of a history of primary headache was greater in patients with diabetes or hypertension than asymptomatic hypertensive patients (45 vs. 27%; p < 0.0001).

Table 1. Clinical, biological and angiographic feature of the whole population with myocardial ischemia and patients stratified by the presence/absence of anginal symptoms

<table>
<thead>
<tr>
<th></th>
<th>Study population (n = 1,427)</th>
<th>Asymptomatic patients (n = 509)</th>
<th>Symptomatic patients (n = 918)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Men, n (%)</td>
<td>1,188 (83)</td>
<td>428 (84)</td>
<td>760 (83)</td>
</tr>
<tr>
<td>Age, years</td>
<td>59±10</td>
<td>59±8</td>
<td>60±9</td>
</tr>
<tr>
<td>Common risk factors, n (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hypertension</td>
<td>703 (49)</td>
<td>244 (48)</td>
<td>459 (50)</td>
</tr>
<tr>
<td>Dyslipidemia</td>
<td>900 (63)</td>
<td>310 (61)</td>
<td>590 (64)</td>
</tr>
<tr>
<td>Diabetes</td>
<td>513 (36)</td>
<td>163 (32)</td>
<td>350 (38)</td>
</tr>
<tr>
<td>Ever-smoking</td>
<td>340 (24)</td>
<td>86 (17)</td>
<td>254 (27)</td>
</tr>
<tr>
<td>Family history of cardiovascular disease</td>
<td>704 (49)</td>
<td>244 (48)</td>
<td>460 (50)</td>
</tr>
<tr>
<td>Stress</td>
<td>677 (47)</td>
<td>219 (43)</td>
<td>458 (50)</td>
</tr>
<tr>
<td>History of primary headache</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Headache history</td>
<td>529 (37)</td>
<td>153 (30)</td>
<td>376 (41)</td>
</tr>
<tr>
<td>No history of headache</td>
<td>898 (63)</td>
<td>356 (70)</td>
<td>542 (59)</td>
</tr>
</tbody>
</table>

No differences in age, gender and common cardiovascular risk factors were observed between symptomatic and asymptomatic subjects (table 1). Patients with symptoms during exercise-induced myocardial ischemia had a significantly greater prevalence of primary headache than those without (41 vs. 30%; p < 0.001).

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patients with a history of angina in daily life than in patients without angina (37 vs. 20%; \( p < 0.001 \)). Of the 534 patients that underwent PTCA, 236 (44%) reported anginal symptoms during the procedure and 301 (56%) reported none. Symptomatic patients during PTCA had a greater prevalence of primary headache than asymptomatic patients (28 vs. 45%; \( p < 0.001 \)). Finally, of the 700 patients with previous MI, 635 (89%) had angina during the event, while 217 (34%) did not; in this subgroup, the prevalence of primary headache was higher in the symptomatic patients than in the asymptomatic patients (34 vs. 17%; \( p = 0.005 \)).

**Discussion**

This study on a large CAD population suggests that the presence of primary headache could be a useful marker to identify subjects with susceptibility to pain and that its absence could be correlated with SMI. In our population, 34% of patients with CAD had a history of primary headache and, in particular, patients with anginal symptoms during the exercise-stress test had a higher prevalence of headache history than asymptomatic patients. This association was present also in the subgroups of hypertensive and diabetic patients and was confirmed by a multivariate analysis. Moreover, a history of primary headache was associated with the presence of referred pain during previous acute MI and/or during PTCA procedures. Taken together, our data suggest that symptomatic patients with primary headache were more prone to pain; this population differs from those with SMI and without headache in their reactivity to pain, confirmed to be a generalized hyposensitivity.

Recent evidence has linked migraine, in particular migraine with aura, to a broader spectrum of ischemic vascular disorders including coronary disease [24–26]. Several factors have been related to the absence of symptoms during myocardial ischemia, but the precise mechanisms responsible for the lack of pain have not yet been fully elucidated [1, 7–9]. Several studies have shown that in patients with SMI, the threshold of pain perception and tolerance are significantly higher than in symptomatic patients [8, 27]. Granot et al. [27] demonstrated that the absence of pain in acute MI was associated with attenuated pain perception in response to various stimuli, and enhanced pain scores were reported in the MI patients with pain compared to those without pain. On the other hand, the onset of headache is widely attributed to a state of hypersensitivity to pain [28]. Therefore, in patients with SMI, the pain threshold was higher than in symptomatic patients, indicating altered systemic modulation of the pain threshold in SMI [8, 29]. However, no previous studies have used the presence/absence of headache to evaluate the pain threshold of SMI patients.

The correlation between primary headaches (especially migraine) and ischemic heart disease has been investigated, but results are still inconclusive [19, 29]. A recent study showed that patients with a history of migraine and other types of headache more commonly have a history of anginal symptoms [28, 29]. As confirmation, a higher prevalence of anginal symptoms has been shown in patients with migraine [17].

In patients with CAD, the correlation between the absence of symptoms during exercise-induced myocardial ischemia and a history negative for headache seems to confirm a generalized hyposensitivity to pain.

Recently, the involvement of intracortical inhibitory circuits was proposed for migraine and an important role in the pathophysiology of headache chronification was proposed for the increase in cortical excitability [30].

A possible role of endogenous endorphins in response to pain has been extensively studied; some studies have suggested a correlation between low levels of endogenous endorphins and anginal symptoms [7, 31, 32], and between low endorphins levels and headache [33].

**Fig. 1.** Distribution of primary headache in patient with and without anginal symptoms during PTCA and MI.
But above all, an important role in response to pain seems to be played by inflammatory substances [34], also implicated in the genesis, progression and clinical manifestations of CAD. In fact, some studies show the presence of anti-inflammatory cytokine pattern in patients with SMI [35, 36]. A certain pattern of anti-inflammatory cytokines may be responsible for the lack of symptoms in patients with SMI; in particular, it is known that some inflammatory cytokines can induce the release of pain mediators such as bradykinin and calcitonin gene-related peptide and activate vagal afferent [37].

One of the most recent hypotheses is linked to the fact that these substances induce an increase of nitric oxide (NO) synthase. An increase of NO has been shown in patients with headache, and it seems that high concentrations of NO are responsible for changes in the presynaptic membrane receptor [38]. However, this role of NO is still controversial and the underlying mechanisms are still poorly understood. For this reason, a role could be played by the opioid system: in a study of patients with chronic tension-type headache, lower levels of Met-enkephalins have been highlighted, confirming their role in influencing the modulation of pain [39].

This pathophysiological mechanism might represent a common denominator between the presence/absence of anginal symptoms and of headache. It is possible to speculate, due to the importance of inflammation in the genesis of atherosclerotic plaque, that the endogenous opioids which derive from local immune cells interact with specific receptors on sensory nerves, leading to strong and clinically measurable analgesia [34, 40]. Similarly, these data seem to confirm that individual differences in pain perception play an important role in the genesis of SMI and, more generally, in determining susceptibility to pain.

The clinical implications of our study may be of interest. Indeed, our data confirm the hypothesis of a generalized hyposensitivity to pain in patients with SMI, who are less susceptible, not only to angina, but also to headache. An interesting aspect of our clinical research indicates that, in patients with documented CAD or at a high risk for CAD who experience headache, there is a greater tendency to experience anginal symptoms: this exposes this population to a lower risk of asymptomatic ischemic episodes. Conversely, for those patients at a high risk for CAD and no history of headache, closer clinical observation is required because the higher prevalence of asymptomatic episodes may put them at increased risk for ischemic events or death.

The symptom headache in a population with risk factors for CAD might be somehow ‘reassuring’ because, according to our data, the occurrence of myocardial ischemia in such population is highly likely to manifest with anginal symptoms. On the other hand, the absence of episodes of headache in a population with risk factors for CAD might suggest a general hyposensitivity to pain. We speculate that such population might have an increased tendency to develop SMI.

In conclusion, our data suggest that the presence of headache in patients with known CAD decreases the probability of their having SMI, whereas the absence of headache recommends a close monitoring for those patients with risk factors for CAD, because in these patients, the risk of developing SMI might be increased. More investigations of a specific nature are needed to clarify this relationship.

References


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