Mental Stress-Induced Myocardial Ischemia

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Abstract
Purpose of Review To summarize recent evidence on mental stress-induced myocardial ischemia (MSIMI), its mechanisms, and clinical significance.

Recent Findings MSIMI can occur in patients with normal cardiac stress testing, is only weakly related to severity of coronary artery disease (CAD), and it is often silent. Among patients with CAD, MSIMI is associated with a twofold increased risk of major adverse cardiovascular events compared to those who do not have MSIMI. Certain groups such as young women with myocardial infarction and those with psychological comorbidities are more susceptible to MSIMI. Abnormal microvascular vasoreactivity and inflammation are implicated mechanisms in MSIMI. Increased brain activity in regions that modulate autonomic reactivity to emotional stress and fear is associated with MSIMI.

Summary MSIMI has important prognostic implications in patients with CAD. Stress can no longer be ignored as a risk factor in cardiology care. Clinical trials testing effective strategies to target MSIMI are needed.

Keywords Mental stress · Cardiovascular reactivity · Microvascular endothelial dysfunction · Coronary vasospasm

Introduction
Coronary artery disease (CAD) remains a leading health threat in the USA [1]. Cardiometabolic risk factors such as hypertension, diabetes, and obesity are major contributors to CAD, but there is unexplained heterogeneity in the risk and outcome of CAD. It is increasingly recognized that psychological stress contributes to CAD risk and outcomes [2, 3], and such effects can be especially powerful among high-risk individuals with preexisting CAD [4]. Stressful exposures such as work stress, interpersonal conflict, low socioeconomic status, early life trauma, and caregiver strain play an important role in CAD in both men and women, but women may be more susceptible to these effects. More recently, mental stress and depression in the COVID-19 pandemic era have been linked to increased cardiovascular risk factors, especially in women [5, 6]. Psychosocial stressors contribute to cardiovascular risk factors by influencing health behaviors (i.e., poor dietary choices, lack of exercise, less medication compliance, and follow-up care) [3, 7], but stress also has direct adverse effects on cardiovascular physiology [8]. Substantial prior evidence links mental stress to endothelial dysfunction [9], atheroma progression, ischemia, arrhythmias, and cardiovascular morbidity and mortality [10–12]. An important potential consequence of stress, among patients with pre-existing CAD, is the provocation of myocardial ischemia. This phenomenon, known as mental stress-induced myocardial ischemia (MSIMI), has been recognized for decades, but it is only recently that its prognostic importance has been fully appreciated. In this review, we will summarize recent findings from our laboratory and others that have used mental stress provocation in patients with...
CAD to study the impact of acute mental stress on cardiovascular physiology, myocardial ischemia, and major adverse cardiovascular outcomes (MACE). We will also discuss neurobiological correlates of these stress-induced alterations.

**Stress Physiology**

Acute mental stress-related ischemia, infarction, and arrhythmias have been described during natural disasters and wars [12–15]. Unexpected enhanced emotional states such as intense anger, extreme fear, panic, bereavement, or even heightened excitement can trigger cardiac events or exacerbate angina in susceptible individuals [8•, 16, 17]. Studies have also linked early trauma and posttraumatic stress disorder (PTSD) to increased risk of cardiac events [17–21]. There are multiple mechanisms by which stress can adversely impact normal cardiovascular physiology (Fig. 1) [8•, 22]. This can be studied in the laboratory setting by exposing participants to various mental stress tasks such as anger recall, exposure to personalized traumatic scripts, mental arithmetic, or a stressful speech task. In response to a mental stress challenge, changes in hemodynamics, electrical conduction, vascular (coronary and peripheral) response, and inflammatory biomarkers are measured to assess stress responses (Table 1). Although assessing transient responses to mental stress remains a laboratory research technique and is not currently translated into clinical practice, this method provides a window as to what may be happening during repeated and chronic daily life stressors in a susceptible individual. Mechanisms of stress physiology in CAD have been recently published [8•]. Here, we summarize key concepts related to stress physiology, focusing on hemodynamic reactivity, vascular function, autonomic function, and inflammation, in patients with CAD [8•].

**Cardiovascular Reactivity (CVR)**

Typically, both heart rate and blood pressure increase in humans exposed to laboratory mental stress, although the increases are greater with exercise stress testing. Both elevated and blunted CVR responses to mental stress have been associated with outcomes [8•, 23]. Greater CVR, defined as changes in blood pressure and heart rate, was associated with myocardial ischemia and was predictive of outcomes in CAD patients [24, 25]. Greater CVR and poor recovery from acute mental stress were associated with an adverse effect on subsequent cardiovascular risk status with higher incident hypertension and increased carotid intima-media thickness [26]. An exaggerated mental stress-induced CVR may trigger acute ischemia in the setting of chronic coronary endothelial dysfunction [27, 28]. Even low CVR is associated with adverse health consequences in some studies [23] and may be a

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**Fig. 1** Mechanisms of stress on cardiovascular physiology. Chronic and acute stressors contribute to an increased inflammatory response and autonomic imbalance, which can lead to adverse cardiometabolic effects, endothelial dysfunction, abnormal vascular reactivity, and abnormal immune function. These factors ultimately can contribute to adverse cardiovascular events. ACTH, adrenocorticotropic hormone; CAM, cell adhesion molecules; E, epinephrine; HTN, hypertension; MMP, matrix metalloproteinase; NAFLD, nonalcoholic fatty liver disease; NE, norepinephrine; RAS, renin-angiotensin system. Figure created using BioRender.com
peripheral marker that signals abnormal neural activity and autonomic dysregulation in certain individuals [23].

Vascular Function

When exposed to acute mental stress, human studies demonstrate both coronary and peripheral endothelial dysfunction [8•, 29–32]. During mental stress, the normal coronary vasodilatory response was found to be blunted in patients with CAD compared to those with normal angiograms [31, 33]. A transient reduction in brachial artery flow-mediated dilation induced by mental stress was associated with adverse cardiovascular outcomes in patients with known CAD during a follow-up period of 3 years [30]. Of note, mental stress responses are highly variable, depending on underlying atherosclerosis and endothelial dysfunction [31–34]. Coronary artery diameter and flow responses to mental stress correlate with coronary diameter responses to acetylcholine, implicating an endothelium-dependent mechanism in abnormal mental stress vasoreactivity [31, 35]. Endothelin-1 (ET-1), a potent vasoconstrictor, has also been implicated in contributing to enhanced vascular reactivity that is induced by sympathetic activation during mental stress stimulus in humans. In healthy participants with no cardiovascular risk factors, selective blockade of endothelin A receptors blocked mental stress-induced peripheral endothelial dysfunction [29]. In one exploratory study, the increase in ET-1 in response to mental stress was associated with parasympathetic withdrawal in CAD patients, but not with circulating catecholamines [36]. In addition to larger brachial artery and epicardial coronary artery reactivity to mental stress, abnormal microvascular reactivity is particularly implicated in mental stress response and discussed in this review.

Table 1  Mental stress myocardial ischemia measures

| Measures of mental stress response | Cardiovascular reactivity | Electrocardiogram and heart rate variability | Cardiac impedance | Macrovascular function | Peripheral microvascular function | Electrodermal activity | Myocardial imaging (nuclear imaging/echo wall motion) | Neuroimaging | Stress biomarkers (blood/urine) |
|-----------------------------------|---------------------------|---------------------------------------------|-------------------|----------------------|-----------------------------------|------------------------|-------------------------------------------------|-------------|-------------------------------|
|                                   | ↑Heart rates and blood pressures during mental stress | ↑Heart rates during mental stress, arrhythmias, ST, and T-wave changes | ↑Heart rate variability during mental stress | ↑Peripheral artery flow-mediated dilation during mental stress | ↑Coronary vasoconstriction during mental stress | ↑Peripheral microvascular vasoconstriction during mental stress | ↑Myocardial ischemia or wall motion abnormality during mental stress | ↑Brain activation during mental stress (e.g., medial prefrontal cortex, anterior cingulate cortex, rostral medial prefrontal cortex, insula, amygdala, hippocampus, hypothalamus, periaqueductal gray) | ↑Circulating biomarkers during mental stress (e.g., cortisol, antidiuretic hormone, epinephrine, norepinephrine, copeptin) |

Autonomic Function

The cardiac autonomic nervous system modulates heart rates, blood pressure, and vascular tone during day-to-day activities and controls physiologic responses to stimuli such as temperature changes, exercise, and stress. Mental stress increases sympathetic nervous system output with accompanying parasympathetic nervous system (vagal) withdrawal. These stress-related changes in autonomic function can be quantified by heart rate variability (HRV), which measures changes in time intervals between heart beats, and are a prognostic marker in cardiac populations [37]. Acute stress-induced sympathetic activation leads to changes in electrophysiology predisposing to abnormal HRV as well as arrhythmias [8•, 22]. Abnormal HRV is also associated with chronic stress factors such as PTSD and depression [8•]. Noninvasive measures such as cardiac impedance-derived pre-ejection period are also used to determine the impact of stress on myocardial physiology (Table 1) [38, 39]. A major area of emerging investigation is the role of neural and autonomic reactivity and triggering of the inflammatory process (see the “Brain Correlates of Angina and Mental Stress Ischemia” section below) [40].

Inflammation

Inflammation is central in CAD pathophysiology that triggers a cascade of events within the arterial wall and leads to development and progression of atherosclerosis [41]. Inflammation and immune dysfunction are key factors in stress response [8•, 42–44]. In addition to catecholamines and stress hormones, mental stress triggers the inflammatory cascade with increased levels of peripheral inflammatory biomarkers, such as interleukin-6 (IL-6) [8•], although timing of the sample collection is important to detect changes
based on the biomarker of interest. Acute stress may trigger plaque rupture through a surge in inflammation but can also activate the coagulation cascade with altered fibrinolysis, resulting in an increased risk for thromboembolism [45, 46]. In a situation of chronic stress, inflammation over time may also predispose to progression of atherosclerosis.

**Mental Stress-Induced Myocardial Ischemia (MSIMI)**

Using cardiac nuclear imaging methods, changes in myocardial blood flow and ischemia can be detected in response to mental stress. Ischemia that is detected with acute mental stress testing done in the laboratory appears to be a phenomenon that is quite distinct from the ischemia that is elicited with exercise or pharmacologic stress testing and is often silent [12, 47, 48]. MSIMI is also linked to factors such as depression as well as to angina and ischemia in daily life (Fig. 2) [24, 49–51]. While MSIMI is not necessarily related to atherosclerotic burden or severity of chest pain, it is emerging as a powerful prognostic marker that is associated with MACE [52••]. In a recent large study by Vaccarino et al. with a total of 918 participants, those who had myocardial ischemia from a standardized mental stress test (a public speech task) were compared to those who did not develop ischemia and also to those who had ischemia from a conventional stress test (exercise or pharmacological) using single-photon emission-computed tomography (SPECT) [52••]. Patients were pooled from two prospective cohort studies: the Mental Stress Ischemia Prognosis Study (MIPS) study and the Myocardial Infarction and Mental Stress Study 2 (MIMS2). The MIPS cohort included patients with CAD, defined based on a previous cardiac catheterization showing atherosclerosis, history of prior myocardial infarction, a history of percutaneous coronary intervention or coronary artery bypass grafting, or a positive nuclear stress test. The MIMS2 cohort included younger patients (ages 25 to 60, 50% women) with a prior history of myocardial infarction within 8 months. The primary endpoint was cardiovascular death or initial or recurrent nonfatal myocardial infarction, and a secondary outcome also included hospitalization for heart failure. In the combined cohort study, 16% of participants developed MSIMI, 31% developed conventional stress ischemia, and 10% were found to have both. MSIMI was significantly associated with elevated cardiovascular death or nonfatal myocardial infarction [52••]. Compared to patients with no

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**Fig. 2** Correlates of mental stress-induced myocardial ischemia (MSIMI). MSIMI is associated with several factors and is a phenomenon that is not benign. MI, myocardial infarction; PTSD, post-traumatic stress disorder.
ischemia, those with MSIMI in the absence of conventional stress ischemia had a twofold increased risk of subsequent adverse events. While conventional stress ischemia in the absence of MSIMI was not significantly associated with adverse outcomes, patients who developed both MSIMI and conventional stress ischemia had an almost fourfold increased risk. These patients also had an increased risk compared to patients with only conventional stress ischemia (Fig. 3).

**Potential Mechanisms of MSIMI**

MSIMI is associated with CAD risk factors, but it is unclear why some individuals with CAD are more susceptible to MSIMI. MSIMI does not appear to be a phenomenon solely of hemodynamic demand triggering ischemia because while standardized mental stress testing typically leads to increases in blood pressure and heart rate, MSIMI occurs at lower levels of oxygen demand compared to that induced by physical stress. In addition to endothelial dysfunction, stress can also cause enhanced smooth muscle reactivity and trigger angina and ischemia episodes [8•, 49, 53–57]. There may be a genetic predisposition to develop MSIMI, as one study has shown a higher likelihood of MSIMI in patients with a β1-adrenergic receptor genetic polymorphism [58]. Although epicardial coronary vasoconstriction can lead to reduced myocardial blood flow, increased microcirculatory tone and resistance induced by mental stress are primarily implicated in MSIMI. Microcirculatory dysfunction (either limited dilation and/or enhanced constriction) is considered a leading mechanism of MSIMI and may trigger myocardial infarction even in the setting of no obstructive CAD, also known as MINOCA. Ersboll et al. reported that patients with MSIMI had left ventricular dysfunction based on impaired resting annular velocities (by echocardiography), even at rest, implicating microvascular dysfunction [59].

Studies have used a peripheral arterial tonometry (PAT) device (Itamar Medical®) to measure peripheral endothelial microvascular vasoconstriction in response to mental stress [32, 60–62]. The device measures changes in microvascular tone with stress as the ratio of pulse wave amplitude during mental stress compared with baseline (prestress). Peripheral vasoconstriction appears to be a correlate of abnormal coronary endothelial function and vasomotor responses [63]. In these studies, the peripheral arterial tonometry ratio during mental stress was consistently lower in patients who developed MSIMI compared with those who did not develop MSIMI, therefore confirming that MSIMI is associated with peripheral vasoconstriction. Furthermore, clear sex differences in hemodynamic and vascular mechanisms associated with MSIMI were found (discussed below) [53, 64]. Circulating norepinephrine (but not epinephrine) levels also predict peripheral vasoconstriction during mental stress [65].

While inflammation increases with mental stress, this increase is not associated with MSIMI. In a recent study, the inflammatory response to mental stress and its relationship

![Fig. 3 Mental stress-induced myocardial ischemia and events in coronary artery disease. MSIMI is associated with twofold increased events in patients with CAD compared to those with no ischemia, and almost a fourfold increased risk when combined with conventional stress ischemia. MI, myocardial infarction. (Figure reprinted with permission from JAMA, 2021, 326(18): 1818–28. Copyright© (2021) American Medical Association. All rights reserved)](image-url)
to MSIMI was investigated in 607 patients (76% men) [66]. As expected, mental stress led to increases in several inflammatory biomarkers such as IL-6, chemoattractant protein-1 (MCP-1), and matrix metalloproteinase 9 (MMP-9) post stress [66]. However, the increase in these biomarkers with stress was not associated with MSIMI [66]. A biomarker of myocardial strain and injury, high-sensitivity cardiac troponin I (hs-cTnI), has also been investigated in relationship to MSIMI. In the MIPS, study levels were measured in 587 patients at rest and after both conventional and mental stress testing. Higher levels of hs-cTnI at rest were associated with MSIMI and conventional stress ischemia. However, there was no significant change in hs-TnI levels post mental stress (at 45 and 90-min post stress), which could be because the laboratory stressor is transient [67]. The association of MSIMI with resting troponin levels suggests that hs-cTnI elevation is an indicator of chronic ischemic burden experienced during everyday life.

**Brain Correlates of Angina and Mental Stress Ischemia**

In addition to cardiac imaging tools and techniques, investigations in recent years have increasingly used brain imaging techniques to understand mechanisms underlying cardiovascular stress physiology and adverse outcomes. In the MIPS study, CAD patients underwent high-resolution positron emission tomography (HR-PET) brain imaging during mental stress tasks [68]. Those with MSIMI compared to those without MSIMI had increased activation with mental stress in the medial prefrontal cortex/anterior cingulate cortex (mPFC/ACC) and related regions [68]. In particular, higher stress-induced activation in the rostromedial prefrontal cortex (rPFC) was associated with MACE at a median follow-up of 3 years, after adjusting for cardiovascular risk factors [40]. MACE was defined as death, myocardial infarction, heart failure hospitalization, and revascularization. The rPFC is an area of the brain that processes stress and regulates autonomic and immune function. The study also found that the association between higher rPFC stress reactivity and adverse outcomes was mediated through vagal withdrawal and systemic inflammatory response pathways [40]. Sex differences were also found in neural responses to mental stress among those with and without MSIMI [69].

When those who reported angina in daily life and those without angina were compared, inferior frontal cortex (IFC) activation during mental stress was independently associated with angina frequency [70, 71]. This region is activated by negative stimuli related to decision-making and social behavior, fear, and emotional stress [71]. There was also increased activation of IFC in patients who had MSIMI [72, 73]. IFC activation is associated with activation in other pain processing regions (thalamus, insula, and amygdala) [70], a network that is activated with negative emotion and interoceptive awareness [71, 74–76]. Higher IFC activity was associated with worsening of angina at 2-year follow-up [70]. Prior work by Rosen et al. also found differences in brain activation patterns during angina in CAD patients compared to those with silent ischemia [77]. In the MIPS study, MSIMI was also found to be associated with cognitive impairment at baseline and with greater decline in cognitive abilities at 2-year follow-up [78].

**Sex Differences**

Based on emerging studies over the past decade that have included more women in mental stress studies, there are notable sex differences in MSIMI. In a sample of patients with stable CAD, Jiang et al. reported that there was a higher prevalence of MSIMI, assessed by echocardiography, in women than in men [79, 80]. In the MIPS study of 686 participants (191 women) with CAD who underwent mental stress testing followed by myocardial perfusion imaging, the incidence of MSIMI in young women (age ≤ 50) was over threefold higher than in age-matched men and older participants [81]. These sex differences in MSIMI were also reported in a prior study, where young women with a history of recent myocardial infarction were found to have more MSIMI detected by nuclear SPECT compared to age-matched men and older participants [82]. In a subsequent larger study of 306 patients (150 women and 156 men) with history of myocardial infarction and a mean age of 50 years, MSIMI was twice as likely in women compared to men [83]. Women in both groups showed greater microvascular constriction to mental stress and a lower reactive hyperemia index with stress. However, there were no sex differences in large artery flow-mediated dilation with mental stress, indicating that a microvascular mechanism is more prominent in women. Indeed, microvascular vasoconstrictive responses predict MSIMI in women [53, 83]. Women, but not men, with MSIMI had more peripheral vasoconstriction compared to their counterparts who did not have MSIMI, whereas men, but not women, with MSIMI had a higher hemodynamic response (measured by rate pressure product change with mental stress) compared to those without MSIMI [53]. These sex differences in MSIMI point to microvascular reactivity being an important mechanism in women, especially since MSIMI was only weakly associated with CAD on coronary angiograms evaluated using the standardized Gensini score [64]. MSIMI was associated with CAD severity in men, but not in women [64].

There are also notable sex differences in the relationship between angina frequency and MSIMI. In the MIPS and MIMS2 samples, angina frequency (in the past month) was
associated with more MSIMI (detected by nuclear imaging) in women but not in men, indicating that angina has important psychological origins among women [49, 84, 85].

It is unclear why young women with CAD are more susceptible to MSIMI. Women have a higher burden of psychosocial risk factors, especially young women, including depression, anxiety, early life adversity, and low socioeconomic status [17, 86, 87]. In the MIMS study, young women with a history of myocardial infarction had more psychosocial risk factors compared to men, although there were no differences in traditional cardiovascular risk factors [83••]. However, even after controlling for these psychosocial factors, the sex differences in MSIMI persisted. It is possible that women’s greater susceptibility towards microvascular dysfunction, which is heightened by sympathetic stimulation during mental stress, plays a role in the higher occurrence of MSIMI among women compared with men.

Inflammation is also actively being investigated as a potential mechanism to explain the differential susceptibility to MSIMI in young women. In a cohort of women (age ≤ 50) with a history of recent myocardial infarction, the levels of IL-6 were two times higher in women compared to men at rest and post mental stress, indicating a higher level of underlying inflammation in women [88]. In the larger combined MIPS and MIMS2 cohort, younger women with CAD had higher levels of inflammation (IL-6) at rest, at 90-min post mental stress, and in response to stress (change in IL-6) compared to young men [46, 88]. These sex differences in inflammatory response were not present in controls without CAD or among older patients. Interestingly, in 562 men and women with stable CAD who underwent a mental stress task, increased inflammation (IL-6 and monocyte chemoattractant protein-1) was associated with MACE (a combined endpoint of death, myocardial infarction, unstable angina with revascularization, and heart failure) among women at 3 years of follow-up, but not among men [42].

Ischemia with No Obstructive Coronary Arteries (INOCA)

Heart disease is the leading cause of death among women in the USA as it is in men, despite that women have less severity of obstructive CAD compared to men. A large proportion of women with angina and ischemia with no obstructive coronary arteries (INOCA) have coronary microvascular dysfunction, which contributes to adverse cardiovascular morbidity and mortality, despite the absence of flow-limiting epicardial stenosis [89–91]. Studies have reported a higher prevalence of anxiety and depression in patients with INOCA compared to those with obstructive CAD [92], although no difference in the prevalence of these factors was found in MI patients with and without obstructive CAD [93]. Given that MSIMI in women is not related to severity of CAD but is related to microvascular responses and to angina frequency, MSIMI could be a potential mechanism to explain chest pain and adverse outcomes in women with INOCA [53, 64].

Women with INOCA show greater peripheral microvascular constriction with mental arithmetic stress compared to age-matched asymptomatic controls [94]. A greater number of participants with INOCA had chest pain during mental stress testing compared to controls (41% vs. 10%, p = 0.01). Higher anxiety and frustration during mental stress correlated with vasodilatation (anxious: r = −0.34, p = 0.04; frustrated: r = −0.37, p = 0.02). While there were no differences in emotional arousal among the groups at baseline (pre-mental arithmetic testing), those with INOCA reported more emotional arousal during the recovery period after mental stress and remained more anxious (p = 0.006), frustrated (p = 0.02), irritated (p = 0.03), and stressed (p = 0.002). Mental stress reactivity, vascular, and autonomic responses in relation to angina burden in women with and without obstructive CAD are under investigation (NCT05401630).

Takotsubo Syndrome

Takotsubo syndrome (TTS) is thought to be an extreme manifestation of MSIMI. In this condition, acute stress exposure triggers a catecholamine storm leading to an exaggerated sympathetic response which in turn can cause acute coronary syndrome and heart failure [16]. In addition to acute increase in left ventricular afterload induced by the catecholamine surge, multivessel coronary vasospasm and coronary microvascular dysfunction contribute to acute myocardial ischemia in this condition [95, 96]. The most characteristic pattern of this syndrome is left ventricular apical stunning with basal hyperkinesis. This syndrome predominantly affects postmenopausal women that have experienced a physical or emotional stressor. In a study of hospitalized patients with TTS, women > 55 years had five times higher odds of TTS compared to women < 55 years old [97]. Approximately 50% of patients with TTS had a history of neurologic or psychiatric disorder compared to 25% of patients with acute coronary syndrome [16]. In a case–control study of patients with whole-body 18-FDG-PET/CT imaging (n = 104, 72% women), higher amygdalar activity was associated with developing TTS after adjustment for risk factors [98].

Potential Treatment

Evidence has mounted that mental stress contributes to myocardial ischemia and adverse outcomes. The next step is to investigate ways to attenuate MSIMI and to determine which strategies translate to improvement in both MACE
and patient-centered outcomes of angina and quality of life. There have been a few studies that have used pharmacotherapy to improve MSIMI [99–101]. In the responses of mental stress induced myocardial ischemia to escitalopram treatment (REMIT) trial, the effects of selective serotonin reuptake inhibitor, escitalopram, were compared to placebo in stable patients with CAD and evidence of MSIMI over a period of 6 weeks [99]. MSIMI was assessed with three different mental stress tasks, and ischemia was detected by echocardiography or electrocardiography during mental stress. Escitalopram therapy led to a reduction in MSIMI (2.6-fold), and interestingly, no difference was noted with exercise-induced ischemia.

Autonomic nervous system modulation with beta-blockers in pilot studies of MSIMI has found variable results [100, 101]. Of note, newer generation vasodilating beta-blockers (i.e., carvedilol and nebivolol) were not tested in these MSIMI studies. In a study of 31 men with CAD who had ambulatory ischemia (using electrocardiographic monitoring) or MSIMI in the lab, those who were randomized to stress management training (1.5-h weekly class) had a lower number of events at 2 years compared to 26 patients randomized to exercise training (three times per week) [102]. This study, however, did not specifically measure the effects of treatment on MSIMI.

Targeting the autonomic nervous system with stress reduction in patients susceptible to MSIMI has not been adequately investigated and may be the next frontier, especially given the role of neural correlates in angina, MSIMI, and MACE. In addition to mind–body interventions and biofeedback techniques [103] that improve autonomic balance, one possible strategy for MSIMI may be vagal nerve stimulation, which was found to reduce sympathetic activity and inflammatory response [38, 104–107].

Conclusions and Future Directions

MSIMI is a unique phenomenon that is different from exercise/pharmacologic stress ischemia and is an independent predictor of adverse cardiovascular prognosis in CAD patients. Evidence thus far indicates that MSIMI is associated with abnormal microvascular reactivity and occurs at lower oxygen demand than conventional stress ischemia. There is substantial variability in responses to acute mental stress, but young women and patients with depression, PTSD, and high anger trait/scores appear to be especially vulnerable to MSIMI. Interdisciplinary mechanistic work combining neuropsychology, cardiovascular stress physiology, and advanced imaging is needed to improve our understanding of the stress pathways that may be playing a critical role in CAD. Research on sex differences in biological responses to psychological stressors (in the acute setting and in everyday life), and on the biological mechanisms that promote resilience, is needed. MSIMI could be used as a treatment target to design effective interventions in appropriate subgroups who may benefit from autonomic modulatory interventions that reduce sympathetic drive. Other important areas of research include assessing whether use of mental stress testing would improve risk stratification in routine clinical care and whether ameliorating MSIMI with interventions would improve outcomes. Even though much work needs to be done, psychological stress must be addressed to improve patients’ care and quality of life. A better integration of mental health and behavioral medicine with routine cardiology clinical care is needed.

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Compliance with Ethical Standards

Conflict of Interest The authors declare no competing interests.

Human and Animal Rights and Informed Consent For the studies conducted by the co-authors that are discussed in this review, informed consent was obtained.

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● Of major importance

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