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Emotional predictors and behavioral triggers of acute coronary syndrome

■ ABSTRACT

Mounting evidence suggests that depression, anxiety, and hostility/anger may each be an independent risk factor for acute coronary syndrome (ACS) occurrence. Data specific to the role of these negative emotional states in predisposing to imminent ACS risk are limited, however. Additionally, a number of studies have indicated that certain situational triggers (such as intense physical exertion) and behavioral triggers (such as acute anxiety or anger) are predictive of imminent occurrence of an ACS. Despite these findings, the use of emotional or behavioral information to identify persons at high risk for imminent ACS onset is not yet practical. Further research is needed to facilitate such patient identification.

Numerous systematic reviews indicate that psychosocial factors are predictive of an initial acute coronary syndrome (ACS),^{1–8} defined as myocardial infarction (MI), unstable angina, or sudden cardiac death. Psychosocial factors under active investigation for their role in ACS onset include chronic negative emotional states (eg, depression, hostility, anxiety) as well as situational and behavioral triggers (eg, acute anger, unusual intense physical activity). This review will present evidence for each grouping of risk markers and then discuss the studies required to better identify and treat patients who have these psychosocial vulnerabilities.

■ NEGATIVE EMOTIONAL STATES

Increasing epidemiologic and pathophysiologic evidence suggests that depression, anxiety, and hostility/anger may each be an independent risk factor for initial occurrence of a cardiovascular event.^{4,8} The vast majority of evidence has been accumulated for depression, and many meta-analyses and systematic

reviews now indicate that depression—as either a clinical diagnosis or an elevation in self-reported symptoms—is a strong, consistent, independent predictor of ACS incidence.^{9–11}

For example, in one meta-analysis of studies examining depression and the incidence of coronary heart disease (CHD) events, the presence of depressive symptoms conferred a relative risk of 1.49 for CHD events (95% confidence interval [CI], 1.16 to 1.92), and the presence of clinical depression conferred a relative risk of 2.69 (95% CI, 1.63 to 4.43), which suggests a dose-response–like association between depression and ACS onset.⁹ **Figure 1** shows the incident risk of CHD events associated with depression (depressed mood and clinical depression) from this meta-analysis relative to the risk associated with established risk factors for CHD.⁴

Major depression

Major depressive disorder is most appropriately diagnosed through direct patient interview, preferably by trained professionals who look for evidence of severely depressed mood lasting at least 2 consecutive weeks, other concomitant symptoms (such as change in eating or sleeping habits), and evidence of associated functional impairment. Because this interview approach is not convenient for large epidemiologic studies, evaluation of major depression by interview in longitudinal epidemiologic studies has been sparse,^{12,13} and only a limited number of other studies have also evaluated the role of a history of depressive disorders¹⁴ or a history of depression treatment.^{14–18}

Depressive symptoms

The impact of self-reported depressive symptoms on cardiac outcomes has been studied more widely—in community settings, among the elderly, and in various cardiac settings—using scales such as the Beck Depression Inventory and the Center for Epidemiological Studies Depression Scale. Notably, these scales assess only the presence of recent depressive symptoms. Nevertheless, despite this important limitation, one-

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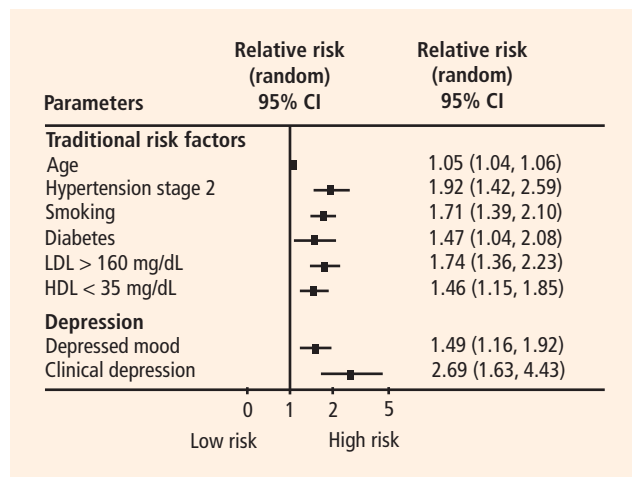


FIGURE 1. Risk of coronary heart disease (CHD) events associated with traditional Framingham risk factors versus the risk associated with depression. Risk ratios for the traditional risk factors are those reported for men followed up in the Framingham study;⁴⁵ risk ratios for depressed mood and clinical depression are from a meta-analysis by Rugulies et al.⁹

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time assessment of healthy participants using these scales has generally revealed a significant relationship between depressive symptoms and future adverse cardiac events.^{9,18} As further evidence of a gradient relationship, the frequency of cardiac events increases as the level of depressive symptoms increases.^{13,19}

Depression is a complex phenotype, and some have argued that subtypes, or intermediary phenotypes, of depression may be particularly predictive of ACS incidence and recurrence.²⁰ Interestingly, the first article ever published documenting a link between depression and coronary artery disease focused exclusively on melancholic depression, or endogenous depression.²¹

Anxiety

Anxiety is defined as a future-oriented negative emotion resulting from perceived threat and accompanied by perceived lack of control and lack of predictability.²² Like depression, anxiety occurs along a continuum, but it is characterized as pathological when it becomes chronic, has intensity out of proportion to any real threat, and leaves the affected person with seriously impaired ability to function.

There are many recognized anxiety disorders, including generalized anxiety disorder, panic attacks, obsessive-compulsive disorder, posttraumatic stress syndrome, and various forms of phobia. As a group, these anxiety disorders constitute one of the most common

forms of psychiatric illness. Moreover, anxiety disorders and major depression are highly comorbid.²³ Although some interesting epidemiologic studies have indicated that anxiety may predict new ACS events,^{24–26} there are a few that have yielded mixed results.^{27,28}

Hostility and anger

Angry and hostile feelings are overlapping emotions and, when experienced frequently, indicate broader, more enduring temperaments or personality styles. Hostility is a cynical, suspicious, and resentful attitude toward others; negative social exchanges, such as sarcasm or impatience, typify individuals with hostility. In contrast, individuals with anger difficulties can have warm, appropriate interpersonal skills and may display verbal aggression or other outbursts only when provoked. Unlike depressive and anxiety disorders, professionally diagnosed, syndromal anger and hostility are not yet recognized by psychiatric nosology.

Relatively few longitudinal studies have been conducted using measures of hostility in healthy cohorts. These studies have reported both the presence^{29,30} and the absence³¹ of positive associations. In a recent systematic review, 7 of 11 studies showed hostility to be a significant risk factor for CHD.⁸ A case-control study involving participants in the Multiple Risk Factor Intervention Trial (MRFIT) employed a structured interview (as opposed to a questionnaire) designed to elicit information regarding signs of hostility, including irritation, arrogance, uncooperativeness, and angry feelings.³⁰ It revealed that men with high hostility levels were more likely to die of cardiovascular disease than men with low hostility levels (adjusted odds ratio = 1.61; 95% CI, 1.09 to 2.39) in this initially healthy but high-risk cohort.³⁰ The investigators chose the structured interview to overcome participants' potential to underreport or underrecognize hostile tendencies when completing a questionnaire.³⁰

Over time, specific measurements of chronic anger,^{32–37} including both unhealthy anger expression³⁴ and suppression,³⁷ have accumulated, permitting examination of their relationship to adverse cardiac outcomes in longitudinal follow-up of disease-free cohorts. A number of these studies have shown a positive association.^{32,33,35–37} Moreover, a recent report from the Framingham study demonstrated an association between anger/hostility and the development of atrial fibrillation and total-cause mortality over a 10-year period,³⁸ and another study has observed a relationship between anger expression and development of stroke.³⁹ Combined, these observations suggest that anger is worthy of further study as a factor in the

TABLE 1**Situational and psychosocial triggers of CHD: Profile of studies assessing their association with imminent ACS***

Trigger	No. of studies	Participants	No. of studies showing association with imminent ACS
Earthquakes	7	Vast majority with elevated risk factors for CVD or clinical histories	6
Sporting events	4	Men	3
War	6	Men	2
Emotional distress and anger	5	Men and women	5
Emotional stress	5	Men and women	5

* Studies extracted from Strike and Steptoe.⁷

CHD = coronary heart disease; ACS = acute coronary syndrome; CVD = cardiovascular disease

development of CHD. As with hostility, however, various subscales have been used to study anger, which makes standardization across studies difficult.

Data specific to imminent ACS risk are few

The majority of the negative emotional states have not been tested for their ability to predict an ACS in the near future. Further, negative emotional and cognitive states that may be implicated in the development of CHD or other cardiovascular disease may be quantitatively or even qualitatively different from those psychosocial factors that identify a patient at imminent risk for an ACS.

SITUATIONAL AND BEHAVIORAL TRIGGERS

Evidence for imminent risk or trigger status can be obtained from retrospective reports, from witness accounts (if available), or prospectively from electronic diaries coupled with ambulatory electrocardiogram recordings or an implantable cardioverter-defibrillator (ICD). Triggers of an ACS include external stimuli (eg, cocaine use, air pollution, ambient temperature), patient activities (eg, eating a meal high in saturated fat, unusual physical exertion), and emotional reactions such as extreme anger or anxiety.^{1,2,7,40,41} Myocardial stunning has also been reported immediately after acute emotional stress but has generally been reversible in these cases.^{42,43} In an observational study, Burg et al found that ICD shocks preceded by an anger episode (as recorded by diary) were more frequent in patients with high trait levels of anger (according to the Spielberger Trait Personality Inventory) and that shocks preceded by an anxiety episode were more frequent in patients with high trait levels of anxiety.⁴⁴ A systematic review by Strike and

Steptoe showed that physical exertion (particularly in poorly conditioned individuals), emotional stress, anger, and extreme excitement are all probable triggers for an ACS.⁷ A recent meta-analysis suggests that emotional stress immediately precedes MI in approximately 7% of MI cases and is a more frequent ACS trigger for women than for men.¹

Table 1 presents examples of triggers that have been found to predict an imminent ACS. Some of these triggers may represent episodes or situations that can lead to an imminent ACS in any apparently healthy person, whereas some may operate only in apparently healthy persons who have chronic negative emotional states or other psychosocial vulnerabilities.

THE PSYCHOSOCIALLY VULNERABLE PATIENT

This selective overview suggests that patients' emotional states are frequently implicated in the onset of an ACS but that the use of behavioral or emotional information to identify those at high risk for imminent ACS onset is not yet practical. Triggers are, by definition, state-like, but their impact may be amplified by trait-like characteristics, such as high dispositional anger, anxiety, hostility, or chronic environmental stress.

The previously mentioned analysis of the MRFIT study by Matthews et al³⁰ suggests that patients identified as "high risk" by conventional risk factors, and who additionally possess high trait hostility, should be monitored closely, as they are at risk for cardiovascular death. Additionally, the Burg study of patients with ICDs suggests that arrhythmia may be induced by acute anxiety or anger in those prone to have such emotions chronically.⁴⁴

Such studies have two research implications. First,

controlled laboratory studies that induce acute negative emotion in those with chronic negative emotional states may help reveal the pathophysiologic processes implicated in immediate ACS onset. Interventions for these psychosocially vulnerable patients await such studies. Second, we must begin to consider that in addition to the psychosocially vulnerable patient, we may have psychosocially vulnerable *situations*, about which we know little at this time.

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