**Paradigm Shift in Angina Management: Integrating Non-Obstructive and Obstructive Causes of Ischemia with Targeted Therapies**

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# Abstract: 150 Words

Management of stable coronary artery disease (CAD) has been based on the plausible assumption that flow-limiting atherosclerotic obstructions are the proximate cause of angina and myocardial ischemia in most patients and represent an important target for revascularization, though the role of revascularization in reducing long-term cardiac events is limited mainly to those with left main disease, 3-vessel disease with diabetes, or decreased ejection fraction. Mounting evidence indicates that non-epicardial coronary causes of angina and ischemia, including coronary microvascular dysfunction, vasospastic disorders, and derangements of myocardial metabolism, are more prevalent than flow-limiting stenoses, raising concerns that many important causes other than epicardial CAD are neither considered nor probed diagnostically. There is a need for a more inclusive management paradigm that uncouples the singular association between epicardial CAD and revascularization and better aligns diagnostic approaches that tailor treatment to the underlying mechanisms and precipitants of angina and ischemia in contemporary clinical practice.

**CONDENSED ABSTRACT: 100 WORDS**

The current management paradigm for coronary artery disease is that flow-limiting atherosclerotic obstructions of epicardial coronary arteries are the proximate cause of angina and ischemia and that coronary revascularization is often the most preferred management. Nevertheless, except for the high-risk subsets with left main disease and low ejection fraction, there is little evidence that revascularization reduces mortality and cardiac events in most other stable angina patients. There is a need for a more inclusive management paradigm that uncouples epicardial coronary obstructions from revascularization and identifies approaches that better align treatment to the underlying mechanisms and precipitants of angina and ischemia.

**KEY WORDS:** Stable angina; myocardial ischemia; coronary microvascular dysfunction; epicardial coronary artery disease, revascularization; percutaneous coronary intervention.

**ABBREVIATIONS LIST**

CABG = coronary artery bypass grafting

CAD = coronary artery disease

CCTA = coronary computed tomography angiography

CMD = coronary microvascular dysfunction

GDMT = guideline-directed medical therapy

ISCHEMIA = International Study of Comparative Health Effectiveness with Medical and Invasive Approaches

INOCA = ischemia and no obstructive coronary arteries

MI = myocardial infarction

PCI = percutaneous coronary intervention

RCT = randomized controlled trial

**Tweet:** There is a need for a paradigm shift in managing stable angina and ischemia. In this article, the authors discuss how both diagnosis and treatment should be tailored to the individual patient, focusing on the many possible pathogenetic causes, including both epicardial coronary obstructions and non-obstructive mechanisms, along with their tailored treatments.**INTRODUCTION: Text Length: 3128 Words Total Word Count: 5065**

Since the advent of coronary angiography more than 60 years ago, stable coronary artery disease (CAD) management has been based on the plausible assumption that "significant" flow-limiting atherosclerotic obstructions of epicardial coronary arteries are the proximate cause of angina and myocardial ischemia in most cases. This belief, supported by anatomic and physiologic evidence that obstructive coronary stenoses can result in regional ischemia and may, in the acute setting, cause acute myocardial infarction (MI), has profoundly influenced our approach to CAD management. In acute MI patients, either percutaneous coronary intervention (PCI) or coronary artery bypass grafting (CABG) can restore coronary flow and improve event-free survival (1, 2). There is also a prevalent belief that epicardial coronary stenosis remains the dominant cause or *sine qua non* of stable angina and ischemia. While, indeed, revascularization may reduce incident cardiac events in high-risk subsets with stable CAD (e.g., left main disease, 3-vessel CAD with diabetes, or decreased ejection fraction), evidence from multiple randomized controlled trials (RCTs) has shown that revascularization of epicardial coronary obstructions, particularly with PCI, does not reduce mortality or morbidity compared with guideline-directed medical therapy (GDMT) in the great majority of stable CAD patients (3, 4).

While revascularization of epicardial stenoses provides better symptom relief and improved quality of life compared to GDMT, recurrence of angina ranges between 20% to 30% within a year after successful PCI (5) and in up to 40% of cases within three years (6), frequently leading to subsequent coronary angiography and repeat PCI. However, since repeat angiography often reveals no evidence of in-stent restenosis or residual coronary obstruction, it is essential to consider non-obstructive causes of angina. Thus, an often-unforeseen consequence of focusing disproportionately on epicardial coronary obstruction is that other pathogenetically important causes of angina and ischemia may not be considered. These causes include epicardial or microvascular coronary vasospasm, coronary microvascular dysfunction (CMD), and derangements of myocardial energy or metabolism(7).

Accordingly, there is a need for a new, more broadly inclusive management paradigm for stable angina patients that uncouples the often-singular association between obstructive CAD and revascularization. Because there are many other potential pathogenetic mechanisms responsible for angina and ischemia, it is essential to identify diagnostic and therapeutic approaches to better tailor appropriate treatment of both obstructive and non-obstructive causes of myocardial ischemia. In so doing, a more pathogenetically-directed approach to diagnosing and treating angina and ischemia would more likely align pharmacologic and procedural interventions as complementary and synergistic for a broader population of stable CAD patients.

# Lessons Learned from Recent Comparative Effectiveness Trials

Earlier RCTs (3,4) showed no incremental benefit of revascularization in reducing mortality, MI, and repeat revascularization when added to GDMT, which included multifaceted pharmacologic secondary prevention and lifestyle intervention. These studies, however, had limitations, e.g., inclusion of low-risk subjects, those with mild-moderate baseline ischemia, use of bare-metal or first-generation drug-eluting stents, and lack of blinding before diagnostic coronary angiography that may have resulted in exclusion of subjects with severe angiographic obstructive disease. The International Study of Comparative Health Effectiveness with Medical and Invasive Approaches (ISCHEMIA) randomized patients with moderate-to-severe inducible ischemia to an initial invasive strategy with revascularization (third-generation drug-eluting stents or CABG) plus GDMT versus an initial conservative strategy of GDMT alone (8). It found no benefit of an invasive approach on the primary endpoint (cardiovascular death, MI, resuscitated sudden cardiac death, hospitalization for unstable angina or heart failure) or secondary endpoint (cardiovascular death or MI). The invasive strategy did result in a statistically significant quality of life improvement, although the overall effect was modest and concentrated mainly in the ~20% of patients with daily/weekly angina (9).

Additionally, a meta-analysis of GDMT with or without PCI in patients with stable CAD (10 RCTs comprising 12,125 patients, including ISCHEMIA) (10) confirmed that PCI did not reduce mortality or MI versus GDMT alone, though the invasive strategy was associated with fewer follow-up revascularizations and improved anginal symptoms.

To evaluate potential bias in unblinded trials, the efficacy of PCI for the treatment of angina was studied in a placebo-controlled trial, which showed no incremental improvement in treadmill walking time, angina relief, or quality of life with PCI + GDMT versus a placebo procedure + GDMT alone (11). While limited by the small sample size and short follow-up, this study raises the issue of whether the observed salutary effect on angina relief attributed to PCI in prior unblinded trials was due, at least in part, to a placebo effect (12).

Finally, we should recognize that managing stable angina patients must include informed, well-considered decision-making involving the patient, family, and physician. Both invasive and conservative approaches may be appropriate and should not be viewed as competing treatment approaches but rather as complementary and potentially additive strategies to enhance optimal patient-centered outcomes (13, 14)

# WHY REVASCULARIZATION MAY Not Be a therapeutic Solution in Many Stable angina Patients

Contrary to type 1 MI for which prompt revascularization is indicated (1,2), revascularization has not been shown to reduce cardiac events in most stable CAD patients (3, 4, 8, 15). Because atherosclerosis is fundamentally a systemic vascular and inflammatory condition affecting epicardial arteries and coronary microcirculation as well as other vascular beds, appropriate GDMT management of ischemia and atherosclerosis must include lifestyle modification (diet, exercise, tobacco cessation), intensive risk factor control and multifaceted pharmacologic secondary prevention (targeting hypertension, dyslipidemia, diabetes, and perhaps inflammation), and when angina is present, effective symptom control (16, 17).

Important observational data from recent, large registries indicate that self-reported angina may improve or resolve over time with medical therapy in most stable CAD patients (18), while subsequent revascularization may be needed only in a minority of patients (~5%) during 5-year follow-up (19). As angina may relapse or remit over time and coronary plaques may become quiescent, an appropriate assessment of angina requires careful follow-up and systematic ascertainment of patient-reported symptoms and quality of life. Thus, a sufficient time horizon (i.e., 3-6 months) is often required for an empiric course of GDMT to be adequately evaluated and efficacy assessed (20, 21). Finally, difficulty in achieving optimal GDMT should not necessarily represent justification to refer patients for revascularization, particularly if a sufficient empiric trial has not been implemented (19-22) or if symptoms are infrequent and mild. Instead, effective GDMT can be achieved by an iterative process that entails collaboration with patients, along with education and counseling, toward a goal of largely patient-directed self-care (23-25).

Nevertheless, ensuring that patients are treated optimally with both lifestyle intervention and multifaceted pharmacologic secondary prevention is time- and labor-intensive, and many cardiologists to whom patients are referred for specific diagnostic testing, including invasive angiography and revascularization, may lack resources to oversee the intensification of medical therapy personally. Thus, more inclusive and coordinated team-management strategies incorporating physician extenders (nurse practitioners, physician assistants, pharmacists) are needed to facilitate optimization of GDMT and improve patient care (23). Using standardized care pathways and management algorithms may further enhance the utilization of these proven approaches (24, 25). Finally, implementation of GDMT likewise remains suboptimal in patients undergoing revascularization (26-28), and such therapies must be similarly prioritized to reduce incident events following revascularization.

**IMPORTANCE OF DIAGNOSING ANGINA AND ISCHEMIA ACCORDING TO THE UNDERLYING PATHOGENETIC CAUSE(S)**

Essential insights on the need for a more encompassing view of the many causes and precipitants of angina and ischemia derive from the SCOT-HEART Trial (29). Here, most patients with known or suspected stable CAD did not have flow-limiting stenoses, indicating that the vast majority (approximately 4 in 5 individuals) had underlying causes of angina and ischemia not attributed to epicardial stenoses (Figure 1). For this reason, a purely anatomical diagnostic approach using invasive coronary angiography or coronary computed tomography angiography (CCTA) may fail to diagnose microvascular and/or vasospastic angina as treatable causes of angina, leading to many patients in whom no obstructive coronary lesions are identified and hence falsely reassured that ischemia is not present. Often such patients are discharged from cardiology, at which point a myriad of potential (and costly) non-cardiac causes are probed rather than pursuing a more diligent evaluation of *non-coronary* causes of angina.

This is particularly important for women, as most patients with ischemia and no obstructive coronary arteries (INOCA) are female (30). Heart disease in women is under-recognized and under-treated, particularly INOCA, where failure to account for microvascular and vasospastic angina within the primarily non-invasive anatomical imaging strategy may result in misdiagnosis (31). Certain stakeholder organizations have recognized that using CCTA as the primary diagnostic testing strategy in angina patients may only help diagnose obstructive epicardial CAD, which is not the most common cause of angina and even less common in women than men.(32).

Indeed, a previous, large observational study of almost 400,000 angina patients undergoing elective coronary angiography found that, among those with a positive non-invasive stress test, only 41% had obstructive CAD (33), indicating a need to embrace a more inclusive management approach that includes many other pathophysiologic mechanisms including CMD and coronary vasospasm (epicardial and/or microvascular) (34–36). Similarly, the 2019 European Society of Cardiology guidelines on chronic coronary syndromes also showed that, among patients with typical angina in the most common age range for detecting stable CAD (50–59 years), 68% of men and 87% of women did not have obstructive coronary stenoses (37), while the Coronary Microvascular Angina (CorMicA) trial (36) and others (34) revealed that approximately 45% of patients presenting with angina or ischemia did not have CAD at angiography. Yet, nearly 90% of these patients demonstrated objective evidence of coronary vasomotor dysfunction (38), including 81% with CMD. Thus, in a sizable proportion of suspected stable CAD patients, CMD or epicardial vasoconstriction can contribute to angina, and because functional mechanisms may co-exist with obstructive CAD, these ischemia precipitants are not necessarily mutually exclusive and may often occur in the same patient (39).

Accordingly, a complete medical evaluation of stable angina patients should characterize the natural history, cardiovascular risk factors, physical examination, and pharmacotherapy (including treatment response, medication intolerance, and adherence). Treadmill exercise testing remains useful to assess functional capacity, the response to the physiological stress of exercise, and limiting symptoms and features of inducible ischemia (notably symptoms and electrocardiographic changes). The response to treatment can be diagnostically informative, and the initial management plan should include antianginal drug therapy, such as short-acting nitrates and either a beta-blocker or a calcium channel blocker. This initial approach complements referral for CCTA since heart rate control (target 60 beats per minute) is required for optimal imaging.

# How to DIAGNOSE AND MANAGE VASOSPASM, MICROVASCULAR dysfunction, and other causes of myocardial ischemia

Both the 2021 AHA/ACC Chest Pain Guideline (40) and the 2019 European Society of Cardiology Chronic Coronary Syndromes Guideline (37) delineate the three different mechanisms of stable angina (obstructive CAD, coronary vasospasm, and coronary microvascular dysfunction). However, a fundamental limitation is the lack of a standard diagnostic evaluation for all patients with suspected angina. While anginal chest discomfort is "the alarm system of the heart" and often the cardinal symptom of myocardial ischemia, it does not provide specificity on its cause. Therefore, it is critical not only to rule in or rule out obstructive CAD but also to establish the cause of myocardial ischemia and to prove or disprove the ischemic origin of symptoms. Such a diagnostic evaluation that comprehensively assesses anatomic and functional coronary alterations would help confirm or exclude the diagnosis of myocardial ischemia and determine the precipitating cause whenever possible.

Myocardial perfusion imaging using positron emission tomography or cardiovascular magnetic resonance imaging is useful for diagnosing CMD (41). These non-invasive imaging techniques provide quantitative and qualitative information on inducible myocardial ischemia. Dynamic first-pass vasodilator stress/rest positron emission tomography uses radiotracers (e.g., 82Rb, 13N-ammonia, 15O-H2O) and quantifies absolute myocardial blood flow (42). Advances with stress cardiovascular magnetic imaging include fully automatic, pixel-wise quantitative mapping of myocardial perfusion (43, 44). This method generates pixel-encoded maps of myocardial blood flow (mL/min/g tissue) during vasodilator stress and at rest. Post-processing software gives accurate measurements for both regional and global stress and resting myocardial blood flow and myocardial perfusion reserve (the ratio of stress to rest myocardial blood flow). A myocardial perfusion reserve < 2.0, in the absence of obstructive CAD, is widely accepted as the CMD threshold associated with adverse outcomes (41).

An algorithm for practical assessment of the multiple causes of angina and ischemia is proposed in the **Central Illustration**. It outlines a pragmatic approach stemming from current international guideline recommendations and results of landmark studies (3, 8, 29). It supports an initial evidence-based approach, including lifestyle interventions and pharmacological secondary prevention with GDMT, to achieve and maintain multiple cardiovascular treatment targets for blood pressure, lipids, and glycemic levels per the current U.S. (40) and European guidelines (37). This algorithm endorses selective functional or anatomic imaging to identify high-risk subsets of stable CAD patients for whom revascularization is more appropriate than medical therapy alone.

If non-invasive studies identify a very low angina threshold and/or a large area of ischemic myocardium at risk during non-invasive stress testing, CCTA or invasive coronary angiography is appropriate to exclude left main and/or high-grade multivessel CAD. In all other chronic stable angina patients, an initial trial of empirical anti-anginal treatment is an important initial step and uptitrating dosages or adding agents for symptom control, as needed, is advocated (21,22). Stable CAD patients with angina should receive at least two antianginal drug classes and adjusted over 3-6-months before referral for revascularization, particularly if anginal symptoms are mild or infrequent. (See Figure 2).

In those with persistent or recurrent ischemic symptoms despite intensive symptomatic treatment, coronary angiography is indicated to identify patients with flow-limiting stenoses who might benefit from myocardial revascularization. In patients without obstructive stenosis, the functional assessment of coronary circulation, including acetylcholine testing for spasm, coronary flow reserve, and microvascular resistance, should be considered to guide subsequent pharmacologic treatment. This algorithm allows tailoring of the diagnostic work up to the clinical situation (**Central Illustration**) and places less emphasis on CCTA which, as currently used, is unable to detect functional coronary alterations (endothelial dysfunction or vasospasm) responsible for ischemia. In SCOT-HEART (29), non-fatal MI at five years was lower in the CCTA-guided group versus the standard care group, but there was no effect on mortality. Secondary prevention therapy, including aspirin and statins, was higher in the CCTA-guided group, further implying that disclosure of atherosclerosis resulted in linked therapy.

Ideally, the above-proposed diagnostic evaluation should be performed in all stable angina patients in whom obstructive CAD has been excluded, but from a practical standpoint, many centers will not have access to such sophisticated testing modalities, or they may lack the skill/expertise to undertake such evaluations, and there are also potential cost-effectiveness concerns that need to be considered. Hence, we advocate additional diagnostic testing, described above, only after obstructive CAD has been excluded and only if symptoms do not improve (or worsen) despite appropriate anti-anginal therapy of at least 2 drug classes (21).

# Why We Need a Paradigm Shift in Our Approach to Angina and Ischemia

Cardiologists should reappraise their thinking of angina and prioritize the following: 1) angina may be due to obstructive CAD and/or INOCA; 2) Most patients presenting with chronic angina do not have epicardial coronary obstructions; 3) if CCTA is the initial diagnostic test and obstructive coronary stenoses are excluded, subsequent testing should include stress perfusion imaging, positron emission tomography, and/or invasive functional coronary angiography with pharmacologic testing to detect coronary microvascular and/or vasospastic mechanisms that may require more targeted therapy; 4) most INOCA patients are women and a diagnostic strategy with a singular focus on defining epicardial coronary obstructions may be inadequate.

Of interest, a comprehensive diagnostic approach that contemplates both anatomical and functional issues can be provided by “dynamic” CCTA and could be viewed as a non-invasive “one-stop shopping” model to diagnose angina and suspected CAD, both obstructive and non-obstructive (45). Ongoing RCTs will determine whether dynamic CCTA fulfills this promise.

**PHARMACOLOGIC MANAGEMENT TARGETING THE PRECIPITANTS OF ANGINA AND ISCHEMIA**

A reduction of coronary flow reserve can cause ischemia due to epicardial stenoses, impaired microvascular function, or both—even in the same patient, as noted above. In this setting, drugs that reduce myocardial oxygen consumption (beta-blockers, non-dihydropyridine calcium channel blockers, or ivabradine) or optimize myocardial oxygen utilization (ranolazine or trimetazidine) are likely the best option. Their combination can also be considered (Figure 2). Alternatively, ischemia can also be caused by epicardial or microvascular spasm. In this setting, vasodilators (calcium channel blockers, nitrates, or nicorandil) are most appropriate, and their combination can also be considered. Thus, to the extent possible, it is highly desirable to tailor pharmacologic therapies to the underlying causes and precipitants of ischemia.

# Conclusion: Where Do We Go from Here?

The time has come for this paradigm shift in managing stable CAD patients. First, we need to expand our current scientific thinking about the many causes and mechanisms of both angina and myocardial ischemia and uncouple the narrow association of ischemia with obstructive epicardial disease (46) as the guiding approach to management. Both angina and ischemia have many causes, but obstructive epicardial disease may or may not be the underlying pathogenetic mechanism (Central Illustration; Figure 2). Hence, our nomenclature should reflect the actual causes of ischemia/angina beyond the currently used terms “coronary” and “disease”, both of which connote epicardial coronary obstruction and are perhaps too narrowly restrictive. A more inclusive and descriptive nomenclature might be considered, such as *“*acute and chronic *myocardial ischemic syndromes”* (47).

Second, we must embrace a more enlightened management approach. Assessments of ischemia that do not delineate abnormal coronary angiographic findings should not necessarily shift diagnostic and therapeutic considerations to *non-cardiac* causes of angina but rather to exploring *non-epicardial coronary* causes (e.g., CMD and vasospastic disorders). We must remain mindful that the evaluation and treatment of angina and ischemia need to be tailored to the individual patient and that adoption of available diagnostic tools required for personalized approaches in clinical practice remains challenging.

Third, we must invest in developing newer management strategies and healthcare delivery models that may better align with treatments proven to benefit patients and society (48–52). Proven secondary prevention strategies and lifestyle interventions that comprise contemporary GDMT continue to be underutilized, particularly in the United States, where as few as 40%–50% of eligible CAD subjects are treated according to established clinical practice guidelines, including those who have been revascularized (26, 49, 51). A recent Viewpoint (52) addressing the new Coronary Artery Revascularization recommendations (53) underscores the critical importance of concomitant preventive therapies in enhancing event-free survival and improving outcomes in stable CAD patients who had undergone CABG or PCI. In this way, perhaps we can re-balance patient management in a way that does not view procedural and pharmacologic interventions as competing treatments but rather as complementary and additive therapeutic approaches best suited to achieving optimal clinical outcomes and symptom relief for our patients.

# HIGHLIGHTS

* There are many pathogenetic mechanisms that may underly myocardial ischemia other than obstructive epicardial coronary disease
* Chronic ischemia management should embrace a more comprehensive evaluation of both epicardial coronary and other causes, with therapy tailored to the underlying precipitants and causes
* A conservative approach to management, including non-invasive testing, lifestyle interventions and goal-directed, multifaceted medical therapy, is evidence-based and effective in many stable angina patients
* Integrating pharmacologic and procedural approaches is essential to optimizing clinical outcomes and should be viewed as both complementary and additive strategies

**Legends to Figures**

**Figure 1. Anatomic Assessment of Myocardial Ischemia and Coronary Artery Disease.**

In SCOT-HEART (33), most patients with suspected stable CAD did not have epicardial coronary obstructions, with the vast majority (approximately 4 in 5 individuals) having angina and ischemia not due to epicardial stenoses.

Abbreviations: CAD = coronary artery disease; CCTA = coronary computed tomography angiography; INOCA = ischemia with no obstructive coronary arteries

**Figure 2. Antianginal Treatment Directed to the Mechanism Responsible for Ischemia**

For exertional angina, anti-anginal drugs that reduce myocardial oxygen consumption (i.e., beta-blockers, non-dihydropyridine calcium channel blockers [CCBs], or ivabradine) are most efficacious whereas for variable threshold angina or CMD, agents that improve myocardial oxygen utilization (i.e., ranolazine or trimetazidine) are suitable treatment options. CCBs are the preferred option for epicardial or microvascular spasm, but nitrates and nicorandil may also be appropriate.

Abbreviations: BB=beta-blockers; BP=blood pressure; CCB=calcium channel blockers; DHP=dihydropyridine; HR=heart rate; LAN=long-acting nitrates

**Central Illustration:** **Management Algorithm for Obstructive and Non-Obstructive Coronary Causes of Angina.** A more inclusive management paradigm for stable CAD patients that addresses the many pathogenetic mechanisms responsible for angina and ischemia is necessary to identify diagnostic and therapeutic approaches that would better tailor the appropriate treatment of obstructive and non-obstructive causes of myocardial ischemia to the underlying ischemia precipitants. Such an approach seeks to promote both evidence-based pharmacologic secondary prevention and procedural interventions as complementary and potentially additive treatments to optimize the management of stable angina patients.

Abbreviations: ACh = acetylcholine; CAD = coronary artery disease; CCTA = coronary computed tomography angiography; CFR = coronary flow reserve; CV = cardiovascular; FFR = fractional flow reserve (a hyperemic pressure ratio); GDMT = Guideline-directed medical therapy; iFR = instantaneous free wave ratio; IMR = index of microvascular resistance; LMD = left main disease

**REFERENCES References Word Count: 1554**

1. Grines C, Patel A, Zijlstra F, et al. Primary coronary angioplasty compared with intravenous thrombolytic therapy for acute myocardial infarction: six-month follow up and analysis of individual patient data from randomized trials. Am Heart J 2003;145:47–57.
2. Kirov H, Caldonazo T, Rahouma M, et al. A systematic review and meta-analysis of percutaneous coronary intervention compared to coronary artery bypass grafting in non-ST-elevation acute coronary syndrome. Sci Rep 2022;12:5138.
3. Boden WE, O’Rourke RA, Teo KK, et al. Optimal medical therapy with or without PCI for stable coronary disease. N Engl J Med 2007;356:1503–16.
4. BARI 2D Study Group, Frye RL, August P, et al. A randomized trial of therapies for type 2 diabetes and coronary artery disease. N Engl J Med 2009;360:2503–15.
5. Ben-Yehuda O, Kazi DS, Bonafede M, et al. Angina and associated healthcare costs following percutaneous coronary intervention: A real-world analysis from a multi-payer database. Catheter Cardiovasc Interv 2016;88:1017–1024.
6. Crea F, Bairey Merz CN, Beltrame JF, et al. Mechanisms and diagnostic evaluation of persistent or recurrent angina following percutaneous coronary revascularization. Eur Heart J 2019;40:2455–2462.
7. Kaski J-C, Crea F, Gersh BJ, Camici PG. Reappraisal of ischemic heart disease. Circulation 2018;138:1463–1480.
8. Maron DJ, Hochman JS, Reynolds HR, et al. Initial invasive or conservative strategy for stable coronary disease. N Engl J Med 2020;382:1395–1407.
9. Spertus JA, Jones PG, Maron DJ, et al. Health-status outcomes with invasive or conservative care in coronary disease. N Engl J Med 2020;382:1408–1419.
10. Shah R, Nayyar M, Le FK, et al. A meta-analysis of optimal medical therapy with or without percutaneous coronary intervention in patients with stable coronary artery disease. Coron Artery Dis 2022;33:91–97.
11. Al-Lamee R, Thompson D, Dehbi H-M, et al. Percutaneous coronary intervention in stable angina (ORBITA): a double-blind, randomised controlled trial. Lancet 2018;391:31–40.
12. Rajkumar CA, Nijjer SS, Cole GD, Al-Lamee R, Francis DP. “Faith healing” and “subtraction anxiety” in unblinded trials of procedures: lessons from DEFER and FAME-2 for end points in the ISCHEMIA Trial. Circ Cardiovasc Qual Outcomes 2018;11:e004665.
13. Ferrari R, Camici PG, Crea F, et al. Expert consensus document: A “diamond” approach to personalized treatment of angina. Nat Rev Cardiol 2018;15:120–132.
14. Yanagawa B, Puskas JD, Bhatt DL, Verma S. The coronary heart team. Curr Opin Cardiol 2017;32:627–632.
15. de Bruyne B, Fearon WF, Pijls NHJ, et al. Fractional flow reserve-guided PCI for stable coronary artery disease. N Engl J Med 2014;371:1208–17.
16. Ridker PM. From CANTOS to CIRT to COLCOT to clinic: Will all atherosclerosis patients soon be treated with combination lipid-lowering and inflammation-inhibiting agents? Circulation 2020;141:787–789.
17. Weintraub WS, Daniels SR, Burke LE, et al. Value of primordial and primary prevention for cardiovascular disease: a policy statement from the American Heart Association. Circulation 2011;124:967–90.
18. Orsini E, Marzilli M, Zito GB, et al. Clinical outcomes of newly diagnosed, stable angina patients managed according to current guidelines. The ARCA (Arca Registry for Chronic Angina) Registry: A prospective, observational, nationwide study. Int J Cardiol 2022;352:9–18.
19. Mesnier J, Ducrocq G, Danchin N, et al. International observational analysis of evolution and outcomes of chronic stable angina: the multinational CLARIFY study. Circulation 2021;144:512–523.
20. Dourado LOC, Poppi NT, Adam EL, et al. The effectiveness of intensive medical treatment in patients initially diagnosed with refractory angina. Int J Cardiol 2015;186:29–31.
21. Boden WE, Kaski JC, Al-Lamee R, Weintraub WS. What constitutes an appropriate empirical trial of antianginal therapy in patients with stable angina before referral for revascularisation? Lancet 2022;399:691–694.
22. Boden WE, Stone PH. To stent or not to stent? Treating angina after ISCHEMIA-why a conservative approach with optimal medical therapy is the preferred initial management strategy for chronic coronary syndromes: insights from the ISCHEMIA trial. Eur Heart J 2021;42:1394–1400.
23. Mason CM. Preventing coronary heart disease and stroke with aggressive statin therapy in older adults using a team management model. J Am Acad Nurse Pract 2009;21:47–53.
24. Brener MI, Tung J, Stant J, et al. An updated healthcare system-wide clinical pathway for managing patients with chest pain and acute coronary syndromes. Crit Pathw Cardiol 2019;18:167–175.
25. Brummel A, Carlson AM. Comprehensive medication management and medication adherence for chronic conditions. J Manag Care Spec Pharm 2016;22:56–62.
26. Xie JX, Gunzburger EC, Kaun L, et al. Medical therapy utilization and long-term outcomes following percutaneous coronary intervention: five-year results from the veterans affairs clinical assessment, reporting, and tracking system program. Circ Cardiovasc Qual Outcomes 2019;12:e005455.
27. Iqbal J, Zhang Y-J, Holmes DR, et al. Optimal medical therapy improves clinical outcomes in patients undergoing revascularization with percutaneous coronary intervention or coronary artery bypass grafting: insights from the Synergy Between Percutaneous Coronary Intervention with TAXUS and Cardiac Surgery (SYNTAX) trial at the 5-year follow-up. Circulation 2015;131:1269–77.
28. Khattab AA, Knecht M, Meier B, et al. Persistence of uncontrolled cardiovascular risk factors in patients treated with percutaneous interventions for stable coronary artery disease not receiving cardiac rehabilitation. Eur J Prev Cardiol 2013;20:743–9.
29. SCOT-HEART investigators. CT coronary angiography in patients with suspected angina due to coronary heart disease (SCOT-HEART): an open-label, parallel-group, multicentre trial. Lancet 2015;385:2383–91.
30. Taqueti VR, di Carli MF. Coronary microvascular disease pathogenic mechanisms and therapeutic options: JACC State-of-the-Art Review. J Am Coll Cardiol 2018;72:2625–2641.
31. Shah NR, Hulten EA, Tandon S, Murthy VL, Dorbala S, Thompson RC. Recent clinical trials support continued emphasis on patient-first over modality-first approaches to initial test selection in patients with stable ischemic heart disease. J Nucl Cardiol 2022 Feb 11. Epub ahead of print.
32. Hobson P, Bakker J. How the heart attack gender gap is costing women’s lives. Br J Card Nurs 2019;14:1–3.
33. Patel MR, Peterson ED, Dai D, et al. Low diagnostic yield of elective coronary angiography. N Engl J Med 2010;362:886–95.
34. Suda A, Takahashi J, Hao K, et al. Coronary functional abnormalities in patients with angina and nonobstructive coronary artery disease. J Am Coll Cardiol 2019;74:2350–2360.
35. Taqueti VR, Hachamovitch R, Murthy VL, et al. Global coronary flow reserve is associated with adverse cardiovascular events independently of luminal angiographic severity and modifies the effect of early revascularization. Circulation 2015;131:19–27.
36. Ford TJ, Stanley B, Sidik N, et al. 1-Year outcomes of angina management guided by invasive coronary function testing (CorMicA). JACC Cardiovasc Interv 2020;13:33–45.
37. Knuuti J, Wijns W, Saraste A, et al. 2019 ESC Guidelines for the diagnosis and management of chronic coronary syndromes. Eur Heart J 2020;41:407–477.
38. Ong P, Camici PG, Beltrame JF, et al. International standardization of diagnostic criteria for microvascular angina. Int J Cardiol 2018;250:16–20.
39. Sechtem U, Brown D, Godo S, Lanza GA, Shimokawa H, Sidik N. Coronary microvascular dysfunction in stable ischaemic heart disease (non-obstructive coronary artery disease and obstructive coronary artery disease). Cardiovasc Res 2020;116:771–786.
40. Writing Committee Members, Gulati M, Levy PD, et al. 2021 AHA/ACC/ASE/CHEST/SAEM/SCCT/SCMR Guideline for the Evaluation and Diagnosis of Chest Pain: A Report of the American College of Cardiology/American Heart Association Joint Committee on Clinical Practice Guidelines. J Am Coll Cardiol 2021;78:e187–e285.
41. Bradley C, Berry C. Definition and epidemiology of coronary microvascular disease. J Nucl Cardiol 2022. May 9. Epub ahead of print.
42. Gupta A, Taqueti VR, van de Hoef TP, et al. Integrated noninvasive physiological assessment of coronary circulatory function and impact on cardiovascular mortality in patients with stable coronary artery disease. Circulation 2017;136:2325–2336.
43. Jacobs M, Benovoy M, Chang L-C, et al. Automated segmental analysis of fully quantitative myocardial blood flow maps by first-pass perfusion cardiovascular magnetic resonance. IEEE Access 2021;9:52796–52811.
44. Kotecha T, Martinez-Naharro A, Boldrini M, et al. Automated pixel-wise quantitative myocardial perfusion mapping by CMR to detect obstructive coronary artery disease and coronary microvascular dysfunction: validation against invasive coronary physiology. JACC Cardiovasc Imaging 2019;12:1958–1969.
45. Serruys PW, Hara H, Garg S, et al. Coronary computed tomographic angiography for complete assessment of coronary artery disease: JACC State-of-the-Art Review. J Am Coll Cardiol 2021;78:713–736.
46. Marzilli M, Merz CNB, Boden WE, et al. Obstructive coronary atherosclerosis and ischemic heart disease: an elusive link! J Am Coll Cardiol 2012;60:951–6.
47. de Caterina R, Boden WE. The nomenclature vagaries for the clinical manifestations of myocardial ischemic syndromes - A call to action. Int J Cardiol 2020;304:5–7.
48. Weintraub WS, Boden WE, Zhang Z, et al. Cost-effectiveness of percutaneous coronary intervention in optimally treated stable coronary patients. Circ Cardiovasc Qual Outcomes 2008;1:12–20.
49. Weintraub WS, Boden WE. Making cardiovascular care more responsive to societal needs. Am J Med 2017;130:1259–1261.
50. Diamond GA, Kaul S. Evidence-based financial incentives for healthcare reform: putting it together. Circ Cardiovasc Qual Outcomes 2009;2:134–40.
51. Bucholz EM, Rodday AM, Kolor K, Khoury MJ, de Ferranti SD. Prevalence and predictors of cholesterol screening, awareness, and statin treatment among US adults with familial hypercholesterolemia or other forms of severe dyslipidemia (1999-2014). Circulation 2018;137:2218–2230.
52. Rhanderson C, Arielle A, E BW, Armin A-Z, Ron B, S BR. The 2021 AHA/ACC/SCAI Coronary Artery Revascularization Recommendations. JACC: Advances 2022;1:1–5.
53. Writing Committee Members, Lawton JS, Tamis-Holland JE, et al. 2021 ACC/AHA/SCAI Guideline for Coronary Artery Revascularization: A Report of the American College of Cardiology/American Heart Association Joint Committee on Clinical Practice Guidelines. J Am Coll Cardiol 2022;79:e21–e129.