The Changing Face of Nuclear Cardiology: Guiding Cardiovascular Care Toward Molecular Medicine

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Radionuclide imaging of myocardial perfusion, function, and viability has been established for decades and remains a robust, evidence-based and broadly available means for clinical workup and therapeutic guidance in ischemic heart disease. Yet, powerful alternative modalities have emerged for this purpose, and their growth has resulted in increasing competition. But the potential of the tracer principle goes beyond the assessment of physiology and function, toward the interrogation of biology and molecular pathways. This is a unique selling point of radionuclide imaging, which has been underestimated in cardiovascular medicine until recently. Now, molecular imaging methods for the detection of myocardial infiltration, device infection, and cardiovascular inflammation are successfully gaining clinical acceptance. This is further strengthened by the symbiotic quest of cardiac imaging and therapy for an increasing implementation of molecule-targeted procedures, in which specific therapeutic interventions require specific diagnostic guidance toward the most suitable candidates. This review will summarize the current advent of clinical cardiovascular molecular imaging and highlight its transformative contribution to the evolution of cardiovascular therapy beyond mechanical interventions and broad blockbuster medication, toward a future of novel, individualized molecule-targeted and molecular imaging-guided therapies.

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T racing its roots to the early 1970s, radionuclide imaging of the heart has been well established as a centerpiece of clinical practice in nuclear medicine (1). Today, nuclear cardiology robustly serves as a gatekeeper to the cardiac catheterization laboratory, by determining the functional and prognostic relevance of coronary artery disease (2). However, in recent years, alternative imaging modalities have emerged for this purpose, backed by evidence from randomized clinical trials—for example, invasive fractional flow reserve as a reference standard for functional guidance of percutaneous coronary intervention (3). Cardiac MRI is as effective as SPECT in reducing the number of invasive coronary angiography procedures (4), and its use as a gatekeeper leads to reduced rates of coronary revascularization compared with assessments with fractional flow reserve (5). Last, the use of coronary CT angiography in patients with stable chest pain lowers the rate of nonfatal myocardial infarction (6), leading to equal outcome when compared with functional imaging, including echocardiography and SPECT (7). This ongoing competition by cardiac MRI, coronary CT angiography, and fractional flow reserve for diagnosis and guidance of intervention in stable coronary artery disease has increased the evolutionary pressure on nuclear cardiology. The resulting refinement of radionuclide imaging methodology has opened avenues for further advances in the field and supports the changing face of nuclear cardiology, in which increasingly specific molecule-targeted imaging techniques find growing clinical applications.

STATE-OF-THE-ART METHODOLOGY FOR CARDIOVASCULAR RADIONUCLIDE IMAGING

Clinical cardiac SPECT is characterized by several methodologic advances. First, there is a strong trend toward shorter acquisition times and reduced radiation exposure. This can be realized using standard dual-head SPECT systems, such as by using innovative focal collimators for cardiocentric acquisitions, along with advanced iterative reconstruction algorithms (8). Alternatively, new solid-state detector technology, such as cadmium-zinc-teLLuride detectors, has been introduced for dedicated cardiac SPECT systems. Acquisition times may be substantially reduced, or radiation dose may be lowered while still achieving image quality superior to conventional SPECT (8).

Beyond SPECT, the use of PET in cardiology is expanding worldwide, supported by its increasing availability due to the success in nuclear oncology (9). Cardiac PET methodology is more accurate than SPECT, and it yields robust absolute quantification (10). Recent developments in ¹¹C-labeled perfusion radiotracers will further increase clinical penetration because of relief from on-site production (11).

Last, nuclear cardiology has also benefited from the widespread acceptance of hybrid imaging technology. Combination of nuclear imaging with CT allows assessment of functional and morphologic abnormalities in a single setting, such as by integrating perfusion assessment with coronary calcium scoring (12). The recent introduction of hybrid PET/MRI adds another dimension, wherein cardiac MRI may provide anatomic, functional, and tissue information while PET adds simultaneous metabolic or molecular information (13). Diagnostic confidence may be also improved by solid-state
silicon photomultiplier digital photon counting PET/CT systems (14). Table 1 summarizes the technical advances of radionuclide cardiac imaging methods.

**RADIOTRACERS FOR CARDIAC IMAGING: FROM FUNCTIONAL TO MOLECULAR IMAGING**

In nuclear cardiology, the major focus in tracer development has traditionally been on identifying the best-suited agent for myocardial perfusion assessment, based on physiologic parameters such as first-pass extraction fraction, proportionality to flow, and tissue retention. These factors remain critical as the field progresses toward absolute flow quantification (15,16). In combination with electrocardiographic gating, function can be better interpreted to guide revascularization (17). Perfusion imaging technically targets physiology and hence is not a molecular imaging technique in its basic sense.

True molecule-targeted imaging nevertheless has a longstanding track record in cardiovascular medicine. Insights into myocardial metabolism are obtained by $^{18}$F-FDG and other radiolabeled substrates, and the modulation of substrate metabolism by ischemia has long been used for PET-based assessment of myocardial viability (18). The spectrum of available molecule-targeted radiopharmaceuticals has expanded, stimulated by advances in other imaging fields. Such pathways include inflammation, fibrosis, and other extracellular matrix processes; angiogenesis; and cell death (Table 2) and have been extensively reviewed elsewhere (19,20).

**FROM MECHANICAL TO MOLECULAR MEDICINE: THE EFFECT OF AVAILABLE THERAPY OPTIONS ON THE EVOLUTION OF CARDIOVASCULAR MOLECULAR IMAGING**

Despite advances in imaging technology and the increasing spectrum of radiopharmaceuticals, until recently molecular imaging in cardiovascular medicine did not reach a level of clinical acceptance similar to that in oncology or neurology. This lack of acceptance is best explained by the fact that targeted molecular therapies in cardiology have also lagged behind those in other fields, and molecular imaging works best in combination with molecular therapies. Traditionally, cardiovascular medicine has been dominated by mechanical interventions. The indication for coronary intervention, valvular intervention, pacemakers, implantable cardioverter-defibrillators, or other devices is effectively established by morphologic or functional imaging, without a clear need for novel molecule-targeted imaging.

The other main pillar of cardiovascular therapy is medical therapy with broadly applicable blockbuster drugs such as angiotensin-converting enzyme inhibitors, β-adrenergic receptor blockers, diuretics, antiplatelet agents, and lipid-lowering agents. Because such drugs show broad benefits and are affordable, their use even for general prevention in a combined poly-pill is under discussion for underserved populations (21). Molecular imaging may identify the target of these drugs directly, such as by visualizing angiotensin-converting enzyme (22) or β-adrenergic receptors (23). But given the low cost of the drugs and their broad applicability, image
TABLE 2
Key Radiopharmaceuticals and Target Mechanisms in Clinical Nuclear Cardiology

<table>
<thead>
<tr>
<th>Tracer</th>
<th>Target mechanism</th>
</tr>
</thead>
<tbody>
<tr>
<td>$^{99m}$Tc-sestamibi, $^{99m}$Tc-tetrofosmin, $^{201}$T</td>
<td>Myocardial perfusion</td>
</tr>
<tr>
<td>$^{99m}$Tc-PYP, $^{99m}$Tc-DPD, $^{99m}$Tc-HMPAO</td>
<td>Cardiac amyloidosis</td>
</tr>
<tr>
<td>$^{123}$I-metaiodobenzylguanidine</td>
<td>Myocardial sympathetic innervation</td>
</tr>
<tr>
<td>$^{123}$I-BMIPP</td>
<td>Fatty acid uptake</td>
</tr>
<tr>
<td>$^{99m}$Tc-annexin V</td>
<td>Apoptotic cell death</td>
</tr>
<tr>
<td>$^{99m}$Tc-RGD imaging peptide</td>
<td>$\alpha_\beta_3$ integrin expression</td>
</tr>
</tbody>
</table>

PET

- $^{15}$O-water, $^{13}$N-ammonia, $^{82}$Rb, $^{18}$F-flurpiridaz*: Myocardial perfusion, absolute myocardial blood flow
- $^{11}$C-deoxyglucose: Glucose utilization (viability), inflammation
- $^{18}$F-sodium fluoride: Microcalcification (atherosclerotic plaque)
- $^{11}$C-PiB, $^{18}$F-florbetaben, $^{18}$F-florbetapir: Cardiac and extracardiac amyloidosis
- $^{11}$C-acetate: Oxidative metabolism, cardiac efficiency
- $^{11}$C-palmitate, $^{11}$C-lactate, $^{11}$C-glucose, $^{18}$F-FTHA: Various aspects of myocardial substrate metabolism
- $^{11}$C-mHED, $^{11}$C-epinephrine, $^{18}$F-LMI1195*: Myocardial sympathetic innervation
- $^{11}$C-CGP12177, $^{11}$C-MQNB, $^{11}$C-KR31173: Receptors of neurohumoral system ($\beta$-adrenergic, muscarinic, angiotensin)
- $^{68}$Ga-pentixafor, $^{68}$Ga-dotatate, $^{11}$C-PK11195: Receptors involved in inflammatory pathways (CXC4, somatostatin type 2, TSPO)
- $^{18}$F-fluciclatide, $^{68}$Ga-FAPI: Targets involved in extracellular matrix reorganization ($\alpha_\beta_3$ integrin, myofibroblast)

$^{18}$F-flurpiridaz and $^{18}$F-LMI1195: preapproval evaluation in clinical trials.

PYP = pyrophosphate; DPD = diphosphonopropane dicarboxylic acid; HMPAO = hydroxymethylene diphosphonate; HMPAO = hexamethyl-propylene-amine-oxime; WBC = white blood cells; BMIPP = $\beta$-methyl-iodophenyl-pentadecanoic acid; RGD = arginyl-glycyl-aspartic acid; PiB = Pittsburgh compound B; FTHA = fluorotheioheptadecanoic acid; mHED = meta-hydroxyephedrine; $^{18}$F-LMI1195 = N-[3-bromo-4-(3-$^{18}$F-fluoro-propoxy)-benzyl]-guanidine; MQNB = methyl-quinclidinyl benzilate; C-XCR4 = CXC-motif chemokine receptor type 4; TSPO = translocator protein; FAPI = fibroblast activation protein inhibitor. $^{11}$C-CGP12177 = $^{11}$C-(4-(3-t-butylamino-2-hydroxypropoxy)-benzimidazol-1); $^{11}$C-MQNB = $^{11}$C-labeled methiodide quinclidinyl benzilate; $^{11}$C-KR31173 = $^{11}$C-labeled 2-Butyl-5-methoxy-methyl-6-(1-oxypropyldin-2-yl)-3-[2-(1H-tetrazol-5-yl)biphenyl-4-yl]methyl]-3H-imidazo[4,5-b]pyridine; $^{11}$C-PK11195 = $^{11}$C-[R]-2-chlorophenyl]-N-methyl-N-[1-methylpropyl]-3-isouquinoline-carboxamide PK11195.

guidance may be ineffective and a clinical role for molecular imaging is unlikely.

As an important development, advanced targeted drugs based on antibodies, small molecules, RNA products, or engineered cells are increasingly introduced into the cardiovascular field, such as for treating inflammatory conditions or interstitial disease. Their specificity may prevent the use of these drugs in broad populations, emphasizing the need for identifying homogeneous patient subsets that benefit most because their underlying disease is driven by a specific mechanism targeted by the respective drug (24). For this purpose, highly specific biomarkers are sought, and molecular imaging–derived biomarkers may have their strongest role (Fig. 1).

**THE SYNERGY OF TARGETED MOLECULAR IMAGING AND THERAPY: LESSONS LEARNED FROM OTHER AREAS OF MEDICINE**

The biomarker reference standard in molecular medicine is an in-depth analysis of tissue obtained from the target region. This, however, requires invasive biopsies that are often difficult to obtain, may suffer from sampling errors, and cannot easily be used for serial follow-up (25). Thus, liquid biopsies have recently been introduced as an alternative; these focus on disease components that can be accessed through analysis of the patient’s blood but neglect intrinsic tumor heterogeneity (25). Imaging biomarkers, on the other hand, do not suffer from the potential risk of sampling error, are noninvasive, provide information about inter- and intraorgan heterogeneity, and allow for noninvasive whole-body read-outs and serial examinations (9). A molecule-targeted radiopharmaceutical assay provides tissue-specific information about the presence or absence of its respective target mechanism and could effectively guide therapy as a virtual biopsy.

In oncology, early evaluation of metabolic response to specific antitumor therapies is conducted with PET in a variety of entities (26). Furthermore, the theranostic principle for neuroendocrine tumors and prostate cancer is rapidly expanding (9). The field of neurology has also been actively pursuing imaging biomarker–guided, targeted therapy concepts, mainly by providing important insights into disease processes followed by novel
Growing role of molecule-targeted therapy in cardiovascular medicine. Targeted therapy is increasingly implemented to support and supplant broad blockbuster drug therapy and mechanical intervention, requiring specific guidance to the right patient at the right time—guidance that may be obtained by molecular imaging.

therapeutic targets. In Parkinson disease, for example, there is a longstanding track record for imaging of the dopaminergic system; such imaging not only may identify disease progression but also may determine the effectiveness of drug therapy (27). In Alzheimer disease, specific tracers targeting pathogenic substrates such as β-amyloid, tau protein, or neuroinflammation have been developed and are increasingly used for guiding and monitoring disease-modifying treatments, which in turn are expected to require improved biomarker guidance for success (28).

Taken together, progress in imaging technology and in radiopharmaceutical development, paralleled by an increased implementation of molecule-targeted therapies in cardiology and supported by proof of feasibility and the success of image-guided molecular therapy in other fields, provides the foundation for the emerging clinical penetration of cardiovascular molecular imaging (Fig. 2).

THE THREE ‘I’S’ (INFECTION, INFLTRATION, AND INFLAMMATION) AT THE VANGUARD OF CLINICAL CARDIOVASCULAR MOLECULAR IMAGING

Cardiovascular device infection, infiltrative cardiomyopathies, and inflammatory conditions of the myocardium and vessel wall are areas that have an increasing clinical need for specific diagnosis and for guidance regarding specific therapy. In these areas, molecular radionuclide–based imaging has been growing rapidly, penetrating the clinical arena.

Infection of Devices

Because of their preeminent survival benefit, implants have seen exceptional success in cardiovascular medicine. Because intracorporeal artificial material supports adherence and growth of bacteria, device recipients are at increased risk of infections associated with substantial morbidity and mortality. Accurate diagnostic tools are needed for early identification of the severity, extent, and location of infection and for guidance of therapeutic options, which may include antibiotic drug therapy, surgical revision, or device extraction (29).

Conventional echocardiography as the first-line test is often limited by device-related artifacts. Because of the potential that electromagnetic fields will interfere with the proper function of cardiac devices, and because of artifacts stemming from the implant material in devices that are safe to operate in a magnetic field, MRI is not an option and CT identifies only morphologic disease aspects (29). Molecule-targeted imaging techniques do not have these limitations. Accordingly, current guidelines propose 18F-FDG PET/CT or white blood cell SPECT/CT in patients with suspected prosthetic valve infection and inconclusive echocardiography findings (29). Of note, PET/CT is not only very sensitive in identifying valvular prosthesis infection but also provides a whole-body assessment for identification of other infectious foci (Fig. 3A). Its diagnostic accuracy is supported by evidence forthcoming from systematic reviews (30), and recent work highlights the importance of early implementation in the diagnostic workup (31). For inconclusive cases, white blood cell SPECT/CT may be of value as a more specific yet time-consuming technique, and cadmium-zinc-telluride technology may further increase accuracy (32).

PET is also expanding toward imaging infection of other devices, including left ventricular assist devices (33). Here, early diagnosis and exact localization of the site of infection may be crucial, because limited involvement of peripheral components is associated with better outcome than involvement of intracardiac components (Fig. 3B) (34). PET effectively guides treatment, which is mostly based on surgical options such as targeted vacuum therapy, substitution of device components, or complete extraction and replacement (33). Molecule-targeted drug therapy options are currently limited to existing antibiotic agents, as novel antiinfective agents are not a focus of industry (35). The potential of molecular imaging to serially monitor antibacterial therapy in device infections has not yet been explored in depth but may represent another area of future growth.

Infiltrative Cardiomyopathies: Amyloidosis

Amyloidosis is a not-so-rare systemic disease characterized by the interstitial deposition of insoluble protein fibrils that affect tissue structure and function. Cardiac involvement may occur in light-chain amyloidosis and in transthyretin-associated amyloidosis (ATTR, either hereditary or acquired) and limits patient prognosis (36). It is established that bone-seeking radiopharmaceuticals identify cardiac involvement in amyloidosis (36). This approach has seen a recent revival driven by 2 major factors: the first is the increasing evidence that cardiac amyloidosis is more prevalent than previously expected, contributing to significant cases of heart failure with preserved ejection fraction (37), and the second is the clinical introduction of novel targeted molecular therapies for amyloidosis—therapies that may be seen as the major contributing factor for the increased use of bone scans. The transthyretin stabilizer tafamidis was approved for cardiac ATTR after the ATTR-ACT trial showed beneficial effects on survival, hospitalization rate, and quality of life (38). In hereditary ATTR, the RNA silencers inotersen and patisiran are additional targeted therapy options that may improve clinical course (39,40), and antisense amyloid protein antibodies may be another promising treatment for various types of amyloidosis (41).

In a recent multisociety consensus document, the usefulness of bone-seeking radiotracers in diagnosing cardiac ATTR was highlighted (42), emphasizing early diagnosis of cardiac involvement in transthyretin gene carriers or subjects with proven
systemic ATTR and new onset of cardiac symptoms. The largest role, however, is in the workup of subjects with newly diagnosed heart failure, for which echocardiography or cardiac MRI show findings suggestive of cardiac amyloidosis. In this setting, a bone scan in the absence of monoclonal gammopathy (which would suggest light-chain amyloidosis) either rules out cardiac amyloidosis if negative or is highly specific for cardiac ATTR in cases of moderate or strong cardiac uptake. Endomyocardial biopsies may be avoided, and therapy decisions can be based on the bone scan, fulfilling the concept of the virtual biopsy (43).

Additionally, PET techniques are emerging in amyloidosis. Although the role of the bone-seeking agent 18F-sodium fluoride for the workup of transthyretin cardiac amyloidosis is still under debate because of low signal intensity (44), PET amyloid markers originally developed for neurology to identify β-amyloid in neurodegenerative disease also detect cardiac amyloidosis. The original 11C-labeled Pittsburgh compound B, as well as the 18F-labeled alternatives florbetaben or flurbetapir, bind to both ATTR and light-chain amyloidosis in the heart, where they allow quantification of cardiac amyloid burden (45). Importantly, amyloid-targeted PET also provides quantitative information about whole-body amyloid distribution (46). A potential area of growth for amyloid PET is in monitoring treatment efficacy in light-chain amyloidosis and ATTR, similar to early response–monitoring approaches in molecular oncology. Ultimately, the available and increasingly used bone-seeking and amyloid-binding molecular imaging agents establish molecular phenotypes of patients (Fig. 4), which will help in early identification of cardiac amyloidosis, guiding patients toward early and effective molecular treatments (43).

Infiltrative or Inflammatory Cardiomyopathies: Sarcoidosis

Sarcoidosis is a systemic granulomatous disease wherein cardiac involvement may occur and contribute to adverse outcome. Like amyloidosis, cardiac sarcoidosis is traditionally underdiagnosed, and this underdiagnosis has been a trigger for increasing implementation of novel, reliable imaging tools. Presently, a combination of 18F-FDG PET (for the assessment of inflammatory disease extent) and resting myocardial perfusion imaging is recommended by expert consensus, wherein thorough dietary preparation is required to suppress myocardial glucose utilization (47). Under the premise of 18F-FDG avidity in granulomatous inflammatory cells, focal or multifocal sites with increased glycolytic activity are indicators of cardiac involvement, particularly with perfusion–metabolism mismatch patterns in which reduced perfusion identifies tissue damage. Nonetheless, focally increased 18F-FDG uptake may not necessarily be specific for sarcoid and has been reported in other inflammatory processes, such as Chagas disease (48). Late-gadolinium-enhancement cardiac MRI studies also accurately identify cardiac sarcoid (49). Yet, caution must be exercised in patients with impaired renal function or implantable devices. Also, PET as a whole-body technique readily identifies extracardiac disease sites (Fig. 5). Most importantly, 18F-FDG PET has the major advantage of specifically identifying inflammatory activity, which predicts outcome and facilitates assessment of therapy response (49). Currently, 18F-FDG PET is considered useful for patients with biopsy-proven extracardiac sarcoid when screening tests (e.g., echocardiography, Holter monitoring, or cardiac MRI) suggest cardiac involvement, unexplained new-onset conduction abnormalities (second- or third-degree atrioventricular block), or idiopathic sustained ventricular tachycardia and when therapy monitoring is needed for proven cases of cardiac sarcoid (47).

Such therapeutic monitoring is of particular potential, because molecular therapeutic options are increasing. Although not limiting the incidence of potentially lethal arrhythmias but, rather, protecting functional myocardium from scar tissue formation (50), corticosteroids remain the first-line therapy in sarcoidosis. Steroid-sparing agents such as antimitabolites (methotrexate, azathioprine, leflunomide) are also embedded in the therapeutic algorithm. In refractory disease, the therapeutic armamentarium has been further expanded by several drugs targeting T-cell activation, Th1/Th17 pathways, or the ubiquitin-proteasome system (51). Thus, as shown for steroids (52), molecular imaging holds promise to address the clinical need to predict and monitor response. It is likely that only patients with active inflammation will respond to antiinflammatory therapies, unlike those with inactive fibrosis or tissue damage. By extension, response to antiinflammatory therapy is best identified by declining inflammatory activity, and when such a response is not detected, an early change in treatment strategy may be pursued (49).

Inflammation of the Vessel Wall

Awareness is growing that inflammatory cascades are key drivers for the onset and progression of atherosclerotic plaques, leading to a higher likelihood of plaque rupture (53). This notion
led to large multicenter trials of antiinflammatory agents. In CANTOS (54), canakinumab, a potent inhibitor of interleukin-1β, reduced the rate of recurrent vascular events in patients with stable coronary artery disease who remain at high proinflammatory risk. In CIRT (55), on the other hand, no benefit from low-dose methotrexate was found in post–myocardial infarction subjects with either diabetes or metabolic syndrome. The discrepant results of these trials emphasize the complexity of inflammation in atherosclerosis. They also indirectly point toward the need for a more precise phenotyping of subjects to select candidates who are most likely to respond. Here, molecular imaging of vessel wall inflammation may have significant potential (56).

After initial work using 18F-FDG (57), various novel radiopharmaceuticals have been explored to visualize plaque biology, including clinically applied agents such as 18F-sodium fluoride as a marker of microcalcification (58), 68Ga-DOTATATE as a marker of macrophage somatostatin receptor expression, and 68Ga-pentixafor as a marker of leukocyte chemokine receptor CXCR4 expression (59). In parallel, the targeting of small vascular structures, especially the coronary arteries, has led to technical refinements of PET imaging by including respiratory and cardiac motion correction (60). These developments supported imaging in selected clinical atherosclerosis trials (61) and generated novel and intriguing hypotheses, as well as tools for future projects on image-guided therapy. A clinical role for molecule-targeted plaque imaging, however, has not yet been established, emphasizing the need for more research into combinations of molecular imaging and therapy in atherosclerosis.

Besides atherosclerosis, other applications for molecular imaging of the vessel wall have emerged. 18F-FDG PET has, for example, entered clinical practice in the diagnosis of large-vessel vasculitis, for which multisociety procedural (62) and clinical use recommendations have been recently published (63). The goal is early diagnosis of vessel wall inflammation indicative of Takayasu or giant cell arteritis and guidance of antiinflammatory therapies. Aortic aneurysm and dissection have also emerged as a target for molecular imaging of microcalcification or inflammatory activity, which may predict progression and complications and guide endovascular repair (64). Finally, similar to cardiac devices and implants, infection of vascular prostheses can also be identified sensitively using 18F-FDG PET (65). The vessel wall with its multiple pathologies therefore remains a challenging but attractive area for future growth of molecular imaging (Fig. 6).

**Inflammation and Myocardial Repair After Ischemic Damage**

Early reperfusion after acute myocardial infarction has led to an improved survival rate, but lower mortality has been accompanied by a rising incidence and prevalence of heart failure (66). Broad blockbuster therapy is used to improve long-term outcome, but this improvement does not change the phenomenon that some patients with acute myocardial infarction completely recover whereas others exhibit progressive contractile dysfunction and left ventricular remodeling. Accordingly, recent efforts have focused on identifying novel therapeutic targets involved in healing and repair after acute myocardial infarction—targets that may be modified to prevent persistent compromise of myocardial function (67). In this regard, postinfarction inflammation has emerged as a key regulator (67), in that an exacerbated inflammatory response to tissue damage may lead to excessive tissue degradation and scar instability, with sustained left ventricular dysfunction (20). Initially, 18F-FDG was advocated to identify the presence of adverse proinflammatory macrophages in subacutely infarcted myocardium (68). Given sufficient metabolic preparation to suppress myocyte 18F-FDG uptake, the metabolic rate of glucose utilization in the infarcted territory correlates with glucose utilization rates in spleen and bone marrow (69), confirming an interrelation between myocardium and lymphoid tissue activation. Subsequently, the 18F-FDG signal in patients early after acute myocardial infarction was confirmed to have a predictive value for later onset of adverse remodeling (70). However, despite thorough preparation for suppression of myocyte uptake, injured but viable myocytes may still show elevated 18F-FDG uptake and thereby compromise the specificity of the signal for inflammatory cells (71). Thus, novel radiotracers with less physiologic accumulation in the myocardium are desirable (20). For example, the translocator protein–ligand 18F-flurticliclamide (GE180) has been investigated in preclinical and clinical settings, confirming utility to obtain further insights into the inflammatory response during wound healing after

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**Figure 3.** 18F-FDG PET/CT of cardiovascular device infection. (A) Prosthetic valve endocarditis: CT, PET, and PET/CT fusion images in representative transaxial and coronal views, along with whole-body maximum-intensity projection (MIP). Intense uptake is present at aortic valve prosthesis, implanted 2 y previously. MIP shows additional focal 18F-FDG–avid embolus in right lower leg (arrow). (B) Left-ventricular assist device infection. Intense uptake is seen around device and outflow tract in left chest wall (left) and along driveline (right) in patient with diffuse reddening and swelling of left chest wall and purulent drainage at entry side of driveline. MIP shows additional uptake in reactive mediastinal lymph nodes.
acute myocardial infarction (72,73). Ultimately, such molecule-targeted imaging approaches may identify individuals with an adverse immune response who may benefit from targeted immunomodulatory interventions to achieve individually optimized myocardial repair (74).

THE FUTURE: EXPANDING OPPORTUNITIES

Innovative Radiopharmaceuticals for Clinical Application: Infection, Inflammation, and Fibrosis

Given the high accumulation of $^{18}$F-FDG in other nonspecific tissue types throughout the body and the inconvenience of preparing radiolabeled white blood cells, novel targets in molecular imaging of infection and inflammation are emerging. Bacteria-specific agents such as maltodextrins, which accumulate specifically in the sites of infection without any interactions, are considered a promising new approach for the diagnosis of device-related infection or other bacterial infectious diseases (75). Additionally, a range of novel agents targeting specific components of the immune system is emerging and holds promise for a more precise assessment of beneficial versus adverse effects of the inflammatory response to tissue injury (75). These agents include, but are not limited to, chemokine receptors such as CXCR4, which has been specifically imaged using the ligand $^{68}$Ga-pentixafor in the myocardium of mice and in patients with acute myocardial infarction (72) and in the atherosclerotic vessel wall (39). Likewise, the $^{68}$Ga-labeled imaging agent DOTA-ECL1i has been introduced for detecting CCR2-positive monocytes and macrophages (76). And last, various mechanisms downstream of early inflammation, which include a complex interplay of reparative, angiogenic, and fibrotic factors, have emerged as imaging targets, because the extent and relative contribution of these processes in response to myocardial injury is thought to be a determinant of myocardial remodeling. These targets include, for example, $\alpha$,$\beta$ integrin as a transmembrane cell surface receptor that interacts with the extracellular matrix and is upregulated after myocardial infarction (75), or activated fibroblasts detected by the novel $^{68}$Ga-labeled fibroblast activation protein inhibitors (77). Ultimately, it is hoped that such innovative imaging compounds will be instrumental not only for more precise diagnosis of individual risk of disease progression but also for personalized guidance of novel targeted therapies (Fig. 7).

Image-Guided Drug Development

Imaging can help in multiple ways to explore, translate, and establish novel targeted therapies in cardiovascular medicine. Although routine morphologic and functional imaging may serve as an endpoint to determine the efficacy of a novel therapy, targeted molecular imaging may be used to characterize the individual disease biology and identify a molecular target that may then be directly manipulated by a specific drug binding via the identical mechanism. This image-guided therapy concept may help in selecting individuals most likely to respond to therapy. Of note, among various novel drugs aiming at modulating the cardiac immune system and fibrosis, specific drugs targeting CXCR4 (78), fibroblast activation protein (77), or other structures amenable for molecular imaging are under evaluation and may be guided by imaging biomarkers. Furthermore, even radionuclide theranostic approaches from nuclear oncology may be translated to cardiovascular medicine. It has, for example, been shown that CXCR4 endoradiotherapy can alter atherosclerotic plaque biology in patients treated for tumors (79), but application in...
primary cardiovascular disease has not yet been pursued. It is highly likely that cardiovascular medicine will see significant growth of such integrated applications of imaging and therapy in the future, opening opportunities for molecular imaging similar to the current success of theranostics in oncology.

Systems-Based Imaging and Therapy: Organ Crosstalk and Cardiooncology

It is increasingly recognized that organ systems do not operate independently, such that injury to one organ can bear grave consequences for disparate organs. Likewise, any targeted drug may exert beneficial effects in a target region but have adverse effects in remote tissue, potentially limiting its overall benefits. Because radiotracers are applied systemically, organ crosstalk and systems-based analyses of networks are readily pursued by whole-body imaging.

Valuable mechanistic insight into organ–organ interactions have been obtained by image-based interrogation of crosstalk between the damaged heart, the hematopoietic system, and the vessel wall (80). More recently, interaction between the brain and the heart in cardiovascular disease has also been elucidated by confirming amygdalar activity as an independent marker of cardiovascular risk (81) and by confirming the presence of neuroinflammation early after myocardial infarction and in chronic heart failure (73). Likewise, interaction between the failing heart and kidneys, lungs, liver, or even gut and microbiome may emerge as a further target for systems-based medical strategies. Molecular imaging of organ interactions will likely be further fueled by the recent introduction of total-body PET/CT, which enables simultaneous coverage and pharmacokinetic studies of the entire human body (82).

A particular area of systemic interaction is the growing field of cardiooncology. Modern tumor therapies are increasingly effective, leading to better survival of cancer patients but introducing adverse effects to organs not directly affected by oncologic disease. Because established and novel antitumor therapies can affect myocardial integrity, systems-based whole-body in vivo imaging may be used to obtain further insights into the off-target response to cancer therapeutics (83). Radiation-, doxorubicin-, or checkpoint inhibitor–induced cardiotoxicity may be identified by

FIGURE 5. $^{18}$F-FDG PET/CT in cardiac sarcoidosis. (A) Representative transaxial CT, PET, and PET/CT fusion images of cardiac region (left) showing strongly increased septal uptake consistent with biopsy-proven active cardiac sarcoidosis. Images of lung region (right), along with whole-body maximum-intensity projection (MIP), show mild, diffuse uptake in both lung lobes indicating moderately active pulmonary sarcoidosis. (B) Integration with cardiac MRI (CMR), showing transmural late gadolinium enhancement in anterior wall, septum, and anterior portion of right ventricle, colocalizing with $^{18}$F-FDG uptake in reangulated short-axis PET/CT images.

FIGURE 6. $^{18}$F-FDG PET/CT of vascular inflammation. (A) Vascular graft infection. CT, PET, and PET/CT fusion images in representative transaxial and coronal views, along with whole-body maximum-intensity projection (MIP), 12 mo after endovascular aortic repair for aneurysm of descending thoracic aorta. CT is consistent with endo-leak of prosthesis, leading to expanding aneurysm, and PET shows intense uptake in periphery of aneurysm, consistent with infection, requiring surgical revision. (B) Large-vessel vasculitis. Shown are representative images in patient with weight loss, elevated level of serum C-reactive protein, and fever of unknown origin. Intense, diffuse elevated uptake in walls of aorta, large arteries of neck, and upper and lower extremities (significantly above liver uptake) is consistent with active giant-cell arteritis.
molecular imaging (83), and assessment of such adverse effects on the cardiovascular system may also offer novel insights into the biology of the myocardium. Thus, given the increasing number of cancer survivors, molecular imaging in cardiooncology is expected to continue to grow.

CONCLUSION

Radionuclide imaging of myocardial perfusion, function, and viability has always been a centerpiece of clinical practice in nuclear cardiology and serves as a reliable tool for therapeutic guidance in ischemic heart disease. However, alternative techniques such as CT or cardiac MRI have applied evolutionary pressure such that nuclear cardiology has evolved from assessing physiologic function and perfusion toward an interrogation of biology and molecular pathways. Today, molecular imaging of the heart is increasingly requested by cardiologists for infection, inflammation, and infiltration in the cardiovascular system. This trend is further fueled by the advent of an increasing spectrum of novel molecule-targeted drug interventions in cardiology, which require early identification of the most suitable patients and subsequent monitoring of success. As such, given the transformative contribution to the evolution of cardiovascular therapy beyond mechanical interventions toward novel, individualized molecule-targeted therapies, the field of molecular cardiac imaging holds strong potential for a future role in precision cardiology.

DISCLOSURE

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