Novel Noninvasive Nuclear Medicine Imaging Techniques for Cardiac Inflammation

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Abstract
Purpose of Review Inflammation is a key player in a wide range of cardiovascular and myocardial diseases. Given the numerous implications of inflammatory processes in disease initiation and progression, functional imaging modalities including positron emission tomography (PET) represent valuable diagnostic, prognostic, and monitoring tools in patient management. Since increased glucose metabolism is a hallmark of inflammation, PET using the radiolabeled glucose analog [18F]-2-deoxy-2-fluoro-d-glucose (FDG) is the mainstay diagnostic test for nuclear imaging of (cardiac) inflammation. Recently, new approaches using more specific tracers to overcome the limited specificity of FDG have emerged.

Recent Findings PET imaging has proven its value in a number of inflammatory conditions of the heart including myocarditis, endocarditis, sarcoidosis, or reactive changes after myocardial infarction. In infection-related endocarditis, FDG-PET and white blood cell scintigraphy have been implemented in current guidelines. FDG-PET is considered as nuclear medical gold standard in myocarditis, pericarditis, or sarcoidosis. Novel strategies, including targeting of somatostatin receptors or C-X-C motif chemokine receptor CXCR4, have shown promising results in first studies.

Summary Nuclear medicine techniques offer valuable information in the assessment of myocardial inflammation. Given the possibility to directly visualize inflammatory activity, they represent useful tools for diagnosis, risk stratification, and therapy monitoring.

Keywords Inflammation · PET · SPECT · Myocardium · FDG · Leukocyte

Introduction

Inflammation contributes to initiation, progression, and healing in various cardiovascular diseases. Its implication has been most thoroughly examined in coronary and vascular atherosclerosis which is now broadly considered an inflammatory disease [1].

Cardiac inflammation can be caused by many different conditions: endocarditis, defined as infection of the endocardial surface of one or more heart valves or of an intracardiac device, has experienced rising incidence in the last decade and is associated with in-hospital mortality rates of 18–23% [2–4]. After myocardial infarction, the balance and orchestration of regional and systemic inflammation plays a crucial role in left ventricular remodeling processes and the subsequent risk of the development of heart failure [5]. Myocarditis, cardiac sarcoidosis, and amyloidosis are under-diagnosed causes of otherwise unexplained cardiomyopathies.

Compared with conventional methods, new noninvasive approaches targeting inflammation have the potential to improve the early detection of myocardial inflammation, enable quantification of disease activity, guide therapeutic interventions, and monitor treatment success. Leukocyte scintigraphy is highly specific for infection because granulocytes are recruited to the site of infection. Whereas general utility has been compromised by limited sensitivity, the implementation of single photon emission computed tomography (SPECT) imaging has increased diagnostic performance and opened
new possibilities in settings in which high specificity is needed, e.g., in endocarditis imaging.

Since increased glucose metabolism is—due to overexpression of glucose transporters and overproduction of glycolytic enzymes in inflammatory cells—considered a hallmark of inflammation, positron emission tomography/computed tomography (PET/CT) using the \(^{18}\text{F}\)-labeled glucose analog 2-deoxy-2-fluoro-d-glucose (FDG) is the standard of reference for molecular imaging of myocardial inflammation. Indications include endocarditis, myocarditis, or sarcoidosis but as well the detection of inflammatory changes after acute myocardial infarction (AMI). However, specificity of FDG is hampered by physiological glucose uptake of the myocardium whose suppression requires dedicated patient preparation. In order to overcome limitations of FDG, a number of promising alternatives have recently been introduced including imaging of somatostatin receptors which are overexpressed on the cell surface of activated macrophages [6]. Furthermore, C-X-C motif chemokine receptor CXCR4, which is also overexpressed by leukocytes, plays a role in stem cell trafficking [7, 8], hence representing a suitable target for molecular imaging. Given the specific nature of their signal, these tracers might be used for noninvasive depiction of the temporal and spatial orchestration of myocardial inflammation. It might thereby be possible to directly assess the extent of inflammatory activity, localize sites of activity, identify therapeutic targets, and monitor response to therapy.

**Myocardial Infarction**

Acute myocardial infarction (AMI) most commonly results from the acute rupture of a coronary atherosclerotic plaque leading to rapid thrombus formation in the infarct-related epicardial artery and subsequent ischemia distal to the site of occlusion [9•]. As a consequence, a well-orchestrated immune response is activated which is crucial for myocardial repair. Overshooting inflammation however has been shown to be associated with inferior outcomes [10, 11].

Cardiac magnetic resonance imaging (CMR) in the clinical setting of AMI can provide information about location and extent of acute myocardial injury as well as left ventricular ejection fraction and end-systolic volume index as important prognostic indicators after AMI [12]. However, no reliable information on inflammatory activity can be obtained. Because of overexpression of glucose transporters and overproduction of glycolytic enzymes in inflammatory cells, PET imaging with FDG is a useful tool in assessing the level of post-AMI inflammation. In experimental models of AMI, FDG-PET has been shown to localize infiltration of metabolically active leukocytes [13, 14•]. Clinical studies supported these findings, providing the opportunity to quantify and monitor metabolic activity in the myocardium after infarction and to gain prognostic information on patient outcome [15, 16•]. The suppression of physiologic FDG-uptake into cardiomyocytes is however mandatory to image monocytes or macrophages and requires patient preparation including low-glucose diet, fasting, and administration of heparin [17, 18]. As potential alternatives to FDG, several novel tracers have been investigated in recent studies: the radiolabeled amino acid \(^{11}\text{C}\)-methionine, routinely used for the characterization of brain tumors and other malignancies, could be demonstrated to serially visualize leukocyte infiltration of inflamed myocardial tissue [19]. Since activated macrophages overexpress somatostatin receptors on the cell surface, the suitability of PET imaging of inflammatory activity using \(^{68}\text{Ga}\)-labeled somatostatin analogs like \(^{68}\text{Ga}\)-DOTA-TATE or \(^{68}\text{Ga}\)-DOTA-TOC has been investigated in a small clinical pilot trial. In comparison to CMR, somatostatin receptor-directed imaging showed a close spatial relation of macrophage concentration to CMR-detected structural changes and might therefore serve as a more specific marker of macrophage activity [20]. However, pre-clinical experiments in mice questioned the usefulness of this approach [21]. Targeting macrophages is also possible by using radiolabeled mannose, an isomer of glucose that is taken up by macrophages through glucose transporters. Additionally, mannose receptors have been described to be expressed on a subset of macrophages [22, 23]. Feasibility of this approach has been shown in atherosclerosis imaging [24•] and may be transferrable also to other settings like AMI.

Another recent approach includes imaging of the C-X-C motif chemokine receptor 4 (CXCR4)/stromal cell-derived factor (SDF) -1α axis which has been shown to play a pivotal role in the recruitment and homing of stem- and progenitor cells to the infarct zone [25, 26]. Local up-regulation of CXCR4 in concert with SDF-1α expression after AMI has been demonstrated to be related to stem cell homing and beneficial outcomes [27–30]. Correspondingly, single systemic treatment with AMD3100, a CXCR4 antagonist, has been shown to result in prolonged bone marrow progenitor cell mobilization and consequently improved recovery from ischemia/reperfusion injury [31]. Additionally, CXCR4 has been described to be routinely expressed on various immune cells and might therefore be involved in the orchestration of the balance between post-infarct inflammation and its resolution [32, 33].

Hence, individual CXCR4 expression after AMI might serve as both a prognostic marker as well as a therapeutic target. Since the advent of a radiolabeled CXCR4-ligand for PET imaging [34] and proof-of-concept for visualization of CXCR4-expression in oncology patients [35–38], some small studies have tried to investigate the translation
of CXCR4-imaging to AMI: local up-regulation of the receptor in the inflamed myocardium as well as in bone marrow and spleen could be demonstrated [39, 40]. Animal models suggested that the imaging signal was mostly created by inflammatory infiltrates as opposed to recruited stem cells (Fig. 1; [40]). Further research to elucidate the potential prognostic and theranostic value of CXCR4 imaging is warranted though.

Information on underlying pathways that might also be targetable for therapeutic interventions might be obtained by imaging P-selectin using [68Ga]-labeled fucoidan, a polysaccharidic ligand of P-selectin with a nanomolar affinity. General feasibility of this approach has been described for vulnerable plaque imaging but might also be transferable to the setting of AMI [41]. Alternative approaches try to image matrix metalloproteinases that are critical in the pathogenesis of disease [16•, 42] or increased expression of αVβ3 integrin as a marker of cardiac repair using the novel PET probe [18F]-Fluciclatide [43].

**Endocarditis**

Early diagnosis of infective endocarditis (IE) remains challenging. Essentially, IE should be suspected in all patients presenting with fever of unknown origin, particularly if it is associated with laboratory signs of infection, anemia, microscopic hematuria, or septic embolic manifestations. The Modified Duke Criteria as the current gold standard include clinical, microbiological, and echocardiographic findings and have proven an overall sensitivity of about 80% [44•]. However, limitations, especially in patients with prosthetic valves (PV) and implantable cardiac electronic devices (ICED) have been reported [45] and can sum up to 24% of...
misclassifications [44•]. Advanced imaging techniques for early and sensitive diagnosis of IE are therefore valuable tools in clinical practice. Combining FDG-PET/CT with the Modified Duke Criteria resulted in increased sensitivity with no major change in specificity [46••]. Whereas FDG-PET is unreliable in the setting of native valve endocarditis [47••], it accurately diagnoses prosthetic valve endocarditis and systemic complications of IE [47••–54•]. Acknowledging this utility, in 2015 FDG-PET/CT was included in the guidelines of the European Society of Cardiology as a major criterion for diagnosing IE in patients with prosthetic valves [55•]. An option to further improve FDG-PET imaging is the incorporation of CT angiography into the PET/CT scan, resulting in 91% sensitivity, 91% specificity, and 93% positive and 88% negative predictive values [53••]. However, specificity of the method may be limited due to artifacts from metal implants or due to the non-specific biologic tracer signal [56, 57]. As a more specific alternative to FDG-PET/CT, the ESC guidelines included SPECT/CT imaging with radiolabeled autologous white blood cells (WBC). Whereas this technique has proven its value in detection of endocarditis [58–60], general application was compromised by limited sensitivity due to a weak signal from the valvular target region and difficult localization of inflammatory foci. The specificity of leukocyte scintigraphy with SPECT/CT could be particularly useful when diagnostic uncertainty remains after echocardiography and FDG-PET/CT, especially in patients who have had cardiac surgery within the past 4 weeks [56, 57, 61, 62].

Additionally, the recent advent of a novel SPECT technology which incorporates a sensitive, heart-focused design with the advantages of cadmium-zinc-telluride (CZT) solid-state detectors seems to infer significant improvements in radionuclide imaging. In an elegant approach combining [111In]-labeled WBC imaging with a simultaneously acquired perfusion study to improve localization of inflammatory hot spots relative to the perfusion-defined valve plane, a German group could demonstrate superior imaging quality as well as increased reader confidence for detection of inflammatory foci (Fig. 2; [63•]).

As another possibility to discriminate between infectious and inflammatory causes of endocarditis, new, more bacteria-specific tracers like carbohydrates exclusively metabolized by bacteria or antibodies directed against components of the bacterial cell membrane, e.g., the pilin protein component of the pilin structure of Enterococcus faecalis are being developed [64].

Cardiac Implantable Electronic Device (CIED) Infections Including Implantable Cardioverter Defibrillator (ICD), Pacemaker-Related and Ventricular Assist Device-Related Infections

Cardiac implantable electronic devices (CIED) have been increasingly used over the last years [55•]. Infection rates are reported to be as high as 1–3% and are associated with a 1-year mortality greater than 10% [65]. Whereas echocardiography is the first line imaging modality for the assessment of supposed CIED infection, its use is severely limited in the investigation of extra-cardiac leads and device pockets. Both FDG-PET/CT and leukocyte scintigraphy with SPECT/CT have proven additional value for diagnosis of ICD- or pacemaker-related infections. FDG-PET/CT has been shown to be especially useful for diagnosis of pocket infection (87–91% sensitivity, 93–100% specificity, 97% PPV, 81% NPV), but is less reliable for diagnosis of lead infection or device-related infective endocarditis (24–100% sensitivity, 79–100% specificity, 66–100% PPV, 73–100% NPV) [66–68]. Nevertheless, in the clinical context of suspected device-related infection, increased and heterogeneous FDG-uptake along a lead appears to be a reliable sign of active infection [66]. Furthermore, presence of a focal hotspot is considered the best criterion of lead infection [69]. Of note, as with all applications involving FDG, accuracy of the examination depends on patient preparation and the interval post-implantation. Mild, unspecific FDG-uptake in patients with an ICD or pacemaker without suspected infection in the acute phase (≤2 months) after cardiac surgery has been reported [57]. Additionally, attenuation artifacts due to metal implants must be avoided by careful scrutiny of non-attenuation corrected images.

Both FDG-PET/CT and leukocyte scintigraphy with SPECT/CT seem to be beneficial in the diagnosis of ventricular assist device (LVAD)-related infection [70, 71]. FDG-PET/CT is especially sensitive for device infection (Fig. 3). In a small retrospective study, sensitivity for LVAD infection was 100% and specificity 80%. Additionally, in 75% of cases, PET imaging had an impact on patient management [72]. Leukocyte scintigraphy with SPECT/CT in LVAD infection has been reported to yield sensitivity, specificity, and positive and negative predictive values of up to 100% each. Moreover, in 23% of cases, otherwise unsuspected extra-cardiac complications could be revealed [71]. The role of FDG-PET-CT in the investigation of extra-cardiac complications of CIED infection has also been investigated. In a retrospective study in patients with suspected CIED infection, whole-body PET also identified distant septic emboli or metastatic infection in 28% of patients [73]. These results could be confirmed in a prospective study in patients with known lead endocarditis [74]. In this cohort, FDG-PET-CT found septic emboli in 10 patients (29%), including 7 cases of spondylodiscitis, 4 of which were not clinically apparent at that point in time and resulted in significant modifications to therapy.

In order to further improve diagnostic accuracy, dual time point protocols for FDG-PET imaging have been proposed. The idea is that imaging at a later time point compensates for background-uptake and might improve accuracy [66, 67, 75]. However, data are limited and studies in other inflammatory
and infectious diseases argue against an added value of this approach [76, 77].

**Myocarditis**

There are many causes for myocarditis, viral infections being the most common. Further etiologies include other types of infections, autoimmune disorders, or drug interactions. The clinical manifestations of myocarditis are highly variable, ranging from subclinical disease to sudden death. The variability in presentation reflects the variability in histological disease severity, etiology, and disease stage at presentation. Myocardial inflammation may be focal or diffuse, involving any or all cardiac chambers. Endomyocardial biopsy (EMB) is currently the gold standard for diagnosing myocarditis with, however, very low sensitivity of only 20–30% and a significant associated procedural risk [78, 79].

CMR is considered the standard imaging modality in the noninvasive diagnosis of myocarditis as it enables detection of various features of myocarditis, including inflammatory hyperemia and edema, myocyte necrosis and scar, changes in ventricular size and geometry, regional and global wall motion abnormalities, and identification of accompanying pericardial effusion [80]. CMR criteria for diagnosis of myocarditis have been summarized as the so-called Lake Louise criteria [81]. However, CMR has its limitations which are particularly apparent in chronic myocarditis with diagnostic accuracies as low as 50% [82]. Additionally, standard CMR may be insensitive for the detection of inflammatory activity, which is critical for monitoring therapeutic responses to prevent secondary tissue alterations.

FDG-PET/CT—after adequate patient preparation—can visualize acute myocardial inflammation to suggest active myocarditis. PET imaging may help to differentiate between active and chronic disease and working protocols have been established [83–85]. In a prospective study enrolling 65 patients with suspected myocarditis, FDG-PET was in good agreement with CMR findings [86]. Given the low yield of random EMB, PET-guided myocardial biopsy may be another application for FDG-PET/CT, as shown for other diseases [87, 88]. CMR and FDG-PET/CT seem complimentary in nature [89] and thus the investigation of the incremental value of integrated PET/MRI for diagnosing myocarditis is an active field of research [84, 90, 91].
In order to overcome limited specificity of FDG, novel PET tracers for imaging of myocarditis are currently investigated. In a rat model of autoimmune myocarditis, feasibility of $[11\text{C}]$-methionine-PET imaging for the detection of cardiac inflammation could recently be demonstrated. Methionine accumulation co-localized with histologically confirmed cardiac inflammatory lesions and $[18\text{F}]$-FDG-uptake, indicating that $[11\text{C}]$-methionine-PET might represent a promising imaging agent for the noninvasive diagnosis of myocarditis (Fig. 4; [92]). Another new approach needing further evaluation includes targeting of somatostatin receptor 2 and has yielded encouraging results in a clinical pilot study [20].

Pericarditis

The causes of pericarditis, acute or chronic inflammation of the pericardium, are manifold and include infection (viral, bacterial, or fungal), myocardial infarction, trauma, malignancy (primary pericardial neoplasms, pericardial metastasis, or paraneoplastic syndrome), autoimmune and inflammatory disease, or metabolic disturbance (uremia). Pericarditis can also be iatrogenic, either postoperative or as a side effect of medication. Radiation therapy or idiopathic causes are other possible origins. Although the etiology is varying, the pericardium has a relatively non-specific response to these different causes: inflammation of the pericardial layers and increased production of pericardial fluids are the most common and often manifest themselves as chest pain.

Echocardiography as the first line of cardiac imaging in the diagnosis or work-up of pericarditis reveals pericardial effusion, pericardial thickening, and septal bounce. Generally, computed tomography and CMR also permit evaluation of pericardial effusion and thickening and permit better differentiation of pericardium and pericardial fluid [93].

The use of FDG-PET/CT in pericarditis is generally complementary and demonstrates its ability to detect inflammatory tissue even in the absence of obvious anatomical changes [94, 95]. Non-infectious and inflammatory pericarditis presents with a mild to moderate FDG-uptake within the pericardium, with either
a diffuse or focal on diffuse pattern of uptake. Only little information is available on the utility of FDG-PET in differential diagnosis of the underlying causes of pericarditis. Some studies reported on the possibility to differentiate infectious/inflammatory pericardial disease from neoplastic/metastatic disease as malignancy often presents with intense metabolic activity [96]. In a retrospective analysis of patients with tuberculous and idiopathic pericarditis, maximum standardized uptake values were significantly higher in tuberculosis patients (13.5 vs. 3.0, \( P < 0.001 \)) [97]. Constrictive or effusive constrictive pericarditis, an uncommon complication of chemotherapy, can present with mild and diffuse pericardial FDG-uptake [94]. Additionally, pericarditis associated with increased metabolism of the wall of the large vessels like the thoracic or abdominal aorta can be caused by underlying vasculitis [98, 99].

### Cardiac Sarcoidosis

Sarcoidosis is a granulomatous disorder of unknown etiology. Most commonly affecting lymph nodes and lungs, it can involve any organ system [100]. Though debated, cardiac involvement is more frequent than previously reported [101, 102] and represents one of the leading causes of death by sarcoidosis in Japan and the USA [103]. Due to the multifocal, patchy pattern of myocardial sarcoid involvement, the sensitivity of endomyocardial biopsy (EMB) is as low as 20–30% [78]. In clinical practice, the Guidelines of the Japanese Ministry of Health and Welfare (JMHWG) which include clinical as well as imaging-based criteria serve as the gold standard for diagnosis of cardiac sarcoidosis (CS) [104]. Though not included in JMHWG, PET/CT using FDG is by far the most commonly used nuclear medicine imaging technique and has mostly replaced \([67\text{Ga}]\)-scintigraphy for assessment of CS [101, 102, 105••, 106].

In comparison to CMR, advantages of FDG-PET (as in other indications) include the biologic nature of the imaging signal, the potential to identify cardiac and extra-cardiac sarcoidosis involvement, and the feasibility of imaging patients with electrical devices or impaired kidney function. Typically, CS manifests as a patchy, focal uptake pattern and FDG-PET/CT has been demonstrated to reliably detect active cardiac and
extra-cardiac sarcoidosis with sensitivities of 81–89% and specificities of 78–82%, respectively (Fig. 5; [106, 107]). FDG-PET is thereby often combined with radionuclide perfusion imaging and electrocardiographic gating in order to rule out coronary artery disease or identify resting perfusion defects suggestive of inflammation-induced tissue damage.

### Classification of Cardiac PET/CT Perfusion and Metabolism Imaging

| Rest Perfusion | FDG | Frequency | Example | Interpretation / Comment |
|----------------|-----|-----------|---------|--------------------------|
| **Normal perfusion and metabolism** | | | | |
| Normal (negative) | Normal | 32 (27%) | | Normal |
| Diffuse (non-specific) | Normal | 15 (12%) | | Diffuse FDG most likely due to failure to suppress FDG from normal myocardium. |
| **Abnormal perfusion or metabolism** | | | | |
| Normal | Focal | 20 (17%) | | Nonspecific pattern; focal increase in FDG may represent early disease vs. normal variant |
| Positive | Negative | 17 (14%) | | Rest perfusion defect may represent scar from cardiac sarcoidosis or other etiologies |
| **Abnormal perfusion and metabolism** | | | | |
| Positive | Focal increase ("mismatch pattern") | 23 (19%) | | Presence of active inflammation ± scar in the same location |
| Positive | Focal on diffuse | 6 (5%) | | Similar to above but also areas of inability to suppress FDG from normal myocardium vs. diffuse inflammation |
| Positive | Focal increase (different area) | 5 (4%) | | Presence of both scar and inflammation but in different segments |

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**Fig. 5** Classification of cardiac PET/CT perfusion and metabolism imaging. Reprinted from [104] by permission of Elsevier/Journal of the American College of Cardiology.
Additionally, FDG-PET/CT in combination with perfusion imaging has proven its value to determine the prognosis of CS patients [105], guiding EMB [88] and in predicting response to and monitoring therapy [110]. In the future, combined FDG-PET/CMR imaging may prove beneficial by combining the strengths of both techniques [111].

Again, physiologic myocardial FDG-uptake requires dedicated patient preparation to avoid the pitfall of inadequately suppressed basal uptake. Protocols include prolonged fasting, dietary modifications (high-fat, low-carbohydrate meals), or application of heparin prior to imaging [17, 112–116]. In order to overcome disadvantages of FDG, more specific radiotracers such as somatostatin receptor-ligands have been investigated in small pilot trials (Fig. 6; [117–119]). Though very preliminary in nature, other alternatives include hypoxia displaying agents like [18F]-fluoromisonidazole (FMISO) [120] or proliferation markers like 3′-deoxy-3′-[18F]-fluorothymidine (FLT) [121].

**Cardiac Amyloidosis**

Cardiac amyloidosis (CA) is a rare form of cardiomyopathy that is characterized by extracellular deposition of fibrils that are composed of low molecular weight subunits of a variety of serum proteins. Although over 25 different amyloidogenic proteins have been described, the three most common types of amyloidosis (defined by their precursor proteins) are light-chain (AL), familial or senile (ATTR), and secondary (AA) amyloidosis. CA most commonly manifests as heart failure, characterized by dyspnea and edema. In its end stage, it presents itself as restrictive cardiomyopathy with a very poor prognosis. Definite diagnosis requires either amyloid deposits on endomyocardial biopsy or, in patients with appropriate cardiac findings, amyloid deposits on histologic examination of a biopsy from other tissues (e.g., abdominal fat pad, rectum, or kidney).

Echocardiography is the initial noninvasive test of choice to diagnose cardiac amyloidosis but suffers from limited specificity [122]. CMR is sensitive and can provide typical pattern of amyloid cardiomyopathy but is strongly limited in patients with moderate to severe kidney disease. Radionuclide bone scintigraphy with technetium-labeled bisphosphonates has long been anecdotally reported to localize to cardiac amyloid deposits. Radionuclide scintigraphy has been reported to be sensitive and specific for imaging cardiac ATTR amyloid and may identify cardiac ATTR amyloid deposits early in the course of the disease [123, 124]. In a recent multi-center trial including 1217 patients with suspected cardiac amyloidosis, the combination of increased myocardial scintigraphic radiotracer-uptake and the absence of a monoclonal protein in serum or urine had a specificity and positive predictive value for cardiac ATTR amyloidosis of 100% [123]. However, scintigraphy cannot reliably detect other types of CA and cannot be quantitatively used for therapy monitoring.

As a potential PET substitute for bisphosphonate-based tracers, general feasibility of amyloid imaging using [18F]NaF has been described in single case series whereas another report failed to identify increased tracer uptake in increased tracer-uptake consistent with inflammatory changes (b, c; arrows). Additionally, enlarged mediastinal lymph nodes and pulmonary nodular lesions were documented. Suspected sarcoidosis was confirmed by transbronchial biopsy. Ten months after treatment initiation, a response of pulmonary (d; insert—baseline CT) and cardiac involvement (e, f) could be recorded. Reprinted from [118] by permission of Oxford University Press/European Heart Journal

**Fig. 6** Macrophage-directed positron emission tomography/computed tomography (PET/CT) using a somatostatin receptor (SSTR) specific ligand ([186Ga]-DOTA-TOC) in cardiac sarcoidosis. A 54-year-old patient with suspected atypical myocarditis was referred. Coronary artery disease was excluded by coronary angiography. Cardiac magnetic resonance imaging (CMR) revealed acute myocardial damage of the septal and anterior wall in T2 and contrast-enhanced images (a). PET/CT using [68Ga]-DOTA-TOC showed corresponding areas of abnormally increased tracer-uptake consistent with inflammatory changes (b, c; arrows). Additionally, enlarged mediastinal lymph nodes and pulmonary nodular lesions were documented. Suspected sarcoidosis was confirmed by transbronchial biopsy. Ten months after treatment initiation, a response of pulmonary (d; insert—baseline CT) and cardiac involvement (e, f) could be recorded. Reprinted from [118] by permission of Oxford University Press/European Heart Journal
ATTR patients [125–127]. Further studies are needed to further investigate the potential value of \(^{18}\text{F}\)NaF PET in CA. Little data available have demonstrated a rather limited role for FDG PET in imaging of CA [128, 129]. To date, the most promising alternatives include more amyloid specific tracers like \(^{11}\text{C}\)-labeled Pittsburgh B (PiB) compound [130, 131] as well as \(^{18}\text{F}\)-labeled compounds such as Florbetapir [132, 133] and Florbetaben [134]. All studies reported on promising results in diagnosis of CA. Furthermore, since lower myocardial uptake of \(^{11}\text{C}\)PiB in AL amyloidosis patients who had recently undergone chemotherapy for monoclonal gammopathy when compared to patients with CA but no prior chemotherapy could be observed, amyloid-directed PET might also be used in therapy monitoring as a surrogate marker of active light chain deposition in the myocardium [130].

**Conclusion**

Myocardial inflammation is the endpoint of various different diseases and pathologic conditions. With FDG-PET/CT and leukocyte scintigraphy being the techniques most commonly used, nuclear medical imaging techniques are powerful tools in diagnosis, risk stratification, and therapy monitoring. Novel approaches including bacteria-specific imaging agents, targeting of somatostatin receptors, or C-X-C motif chemokine receptor CXCR4 can help to overcome limitations of standard functional imaging techniques and have demonstrated encouraging results in pilot studies.

**Compliance with Ethical Standards**

**Conflict of Interest** MK and CL declare that they have no conflicts of interest.

**Human and Animal Rights and Informed Consent** All reported studies/experiments with human or animal subjects performed by the authors have been previously published and were in compliance with all applicable ethical standards (including the Helsinki declaration and its amendments, institutional/national research committee standards, and international/national/institutional guidelines).

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