Imaging cardiac innervation in amyloidosis
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Cardiac amyloidosis is a form of restrictive cardiomyopathy resulting in heart failure and potential risk on arrhythmia, due to amyloid infiltration of the nerve conduction system and the myocardial tissue. The prognosis in this progressive disease is poor, probably due the development of cardiac arrhythmias. Early detection of cardiac sympathetic innervation disturbances has become of major clinical interest, because its occurrence and severity limits the choice of treatment. The use of iodine-123 labelled metaiodobenzylguanidine ([I-123]MIBG), a chemical modified analogue of norepinephrine, is well established in patients with heart failure and plays an important role in evaluation of sympathetic innervation in cardiac amyloidosis. [I-123]MIBG is stored in vesicles in the sympathetic nerve terminals and is not catabolized like norepinephrine. Decreased heart-to-mediastinum ratios on late planar images and increased wash-out rates indicate cardiac sympathetic denervation and are associated with poor prognosis. Single photon emission computed tomography provides additional information and has advantages for evaluating abnormalities in regional distribution in the myocardium. [I-123]MIBG is mainly useful in patients with hereditary and wild-type ATTR cardiacl amyloidosis, not in AA and AL amyloidosis. The potential role of positron emission tomography for cardiac sympathetic innervation in amyloidosis has not yet been identified. (J Nucl Cardiol 2017)

Key Words: Amyloidosis • Sympathetic • Innervation • MIBG

INTRODUCTION

Patients with amyloidosis are prone to developing disturbances in autonomic innervation: dysautonomia.¹

Cardiac dysautonomia can be caused by amyloid infiltration into the myocardial and conduction tissue, resulting in conduction and rhythm disorders. Cardiac dysautonomia is common in patients with transthyretin-related amyloidosis (ATTR type) and in patients with immunoglobulin light chain-derived amyloidosis (AL type).² More specific, patients with the hereditary form of ATTR type amyloidosis (hATTR, formerly called familial amyloid polyneuropathy) frequently develop polyneuropathy and dysautonomia. Furthermore, cardiac dysautonomia may occur independent of the presence of a typical restrictive cardiomyopathy. Amyloidosis’ typical restrictive cardiomyopathy is most commonly found in patients with wild-type ATTR type amyloidosis (wtATTR, formerly called senile systemic amyloidosis). In these wtATTR patients, polyneuropathy and dysautonomia are infrequent and approximately 9%.³
At present, actual amyloid infiltration cannot be visualized with nuclear medicine techniques. Nonetheless, semi-quantitative analysis of tracer accumulation in the left ventricle compared to the background (heart-to-mediastinum ratio, HMR) on iodine-123 labelled metaiodobenzylguanidine ([I-123]MIBG) scintigraphy, is assumed to provide insight in the amyloid infiltration of the sympathetic nerve system.4–12 [I-123]MIBG, a chemically modified analogue of norepinephrine, is stored in vesicles in presynaptic sympathetic nerve terminals and not further catabolized. Decreased HMR at 4 h after tracer administration (late HMR) reflects the degree of sympahtethical dystonia, and is found to be an independent prognostic factor in the development of ventricular dysrhythmia.13 Whereas showing promising results in ischemic heart disease, positron emission tomography (PET) for sympathetic innervation in cardiac manifestation of amyloidosis has not yet been studied.14

The purpose of this review is to provide an overview of the present literature on the application of nuclear imaging modalities for the evaluation of cardiac innervation in patients with amyloidosis, and its future perspectives (Figures 1, 2, 3).

**METHODS**

For this review a literature search was performed on PubMed on May 5th 2017, using the following string: "[(innervation) OR (sympathetic)] AND [(amyloidosis) OR (amyloid)] AND [(heart) OR (cardiac)] AND [(nuclear) OR (imaging)]", resulting in 29 hits, of which 24 were considered relevant for this review. Reviews, editorials, abstracts, case reports, animal studies, conference presentations were excluded. In total, 16 articles were found that used radiopharmaceuticals for conventional nuclear medicine imaging of cardiac innervation, all with [I-123]MIBG. The results of these papers are summarized below and divided into three main topics: the imaging of cardiac innervation itself, the implications of this imaging method, and the relation with other nuclear medicine imaging techniques in cardiac amyloidosis.

**IMAGING OF CARDIAC INNERVATION IN AMYLOIDOsis**

Imaging of cardiac innervation in patients with amyloidosis has been mainly focused on visualizing the effects of amyloidosis on the sympathetic nerve system. Conventional nuclear imaging [I-123]MIBG is the most widely used modality for this indication. Table 1 provides an overview of the present available literature with respect to the use of [I-123]MIBG in patients with different types of systemic amyloidosis. The main results regarding HMR, wash-out and patient outcome of the different studies are displayed. As shown in this overview, hATTR type amyloidosis patients are studied most extensively, showing the most pronounced reduced late HMR. Also AL type amyloidosis patients tend to have decreased late HMR compared to healthy control subjects, however to a lesser extent compared to both hATTR and wtATTR type patients.8,10,12 Due to the large overlap of late HMR ranges in ATTR and AL type amyloidosis patients, [I-123]MIBG scintigraphy is considered not to be able to discriminate between these amyloidosis subtypes.12 Despite that cardiac manifestations are very rare in patients with secondary (AA) amyloidosis, one study showed lower mean late HMR in 11 AA type amyloidosis patients compared to healthy control subjects.12 This finding may contribute to the assumption of amyloid deposits infiltrating the conducting system during the course of the disease.

Mean late HMR differs substantially between the different publications. This variability is mainly due to non-homogeneity in [I-123]MIBG imaging acquisition. HMR varies between different gamma camera systems (vendors), but more importantly between the application of low energy and medium energy collimators.15 Generally, HMR is higher on images acquired with medium energy collimators compared to images acquired with low energy collimator.16 Based on these differences in HMR, cut-off values for the different collimators are proposed, as well as conversion algorithms.17,18 Additional single photon emission computed tomography (SPECT) scanning may be of value in the evaluation of regional cardiac sympathetic innervation abnormalities. The majority of patients (both AL and ATTR type amyloidosis) with low HMR show reduced tracer accumulation in the infero-postero-lateral segments.4,6,11 Unfortunately, this may not be considered as a characteristics finding in amyloidosis patients, since a defect in [I-123]MIBG accumulation in the inferior myocardial wall is also reported in healthy control subjects.19 This is considered as a consequence of physiological [I-123]MIBG accumulation in the liver overprojecting the infero-posterior myocardial wall.

**IMPLICATIONS OF IMPAIRED CARDIAC SYMPATHETIC INNERVATION**

Studies using [I-123]MIBG in patients with ischemic heart disease (IHD) have shown that disrupted cardiac sympathetic innervation based on low late HMR is associated with an increased risk on developing ventricular arrhythmia and appropriate implantable cardioverter-defibrillator (ICD) shocks, and is associated with poor survival.13,20 In fact, reduced late HMR is a
stronger prognostic factor than left ventricular ejection fraction (LVEF) for developing severe adverse cardiac events in patients with IHD.\textsuperscript{13} In amyloidosis patients with impaired cardiac sympathetic innervation, decreased survival rates are also established.\textsuperscript{21–23} Late HMR was identified as an independent prognostic factor for 5-year all-cause mortality, with a 42\% mortality rate for those patients with late HMR <1.60, compared to merely 7\% in patients with late HMR \( \geq 1.60 \) (hazard ratio (HR) 7.2, \( P < 0.001 \)).\textsuperscript{21} Based on the results of this study, even patients with HMR <1.60 seem to benefit from liver transplantation (because of amyloid involvement), resulting in lower long-term mortality than neurophysiological score-matched control subjects (HR 0.32, \( P = 0.012 \)).\textsuperscript{21} This underlines the assumption that impaired cardiac sympathetic innervation will not

**Figure 1.** Example of a 70 year old female patient with ATTR amyloidosis based on Val30Met mutation, with both positive bone scan (A) and [I-123]-MIBG scintigraphy. B 15 minutes post injection (p.i.), C 4 hours p.i.. Late HMR 1.38, normal value in our laboratory: 2.0, performed with a medium energy collimator.
progress after liver transplantation, and that re-innerva-
tion cannot be detected within this duration of clinical
follow-up.11,23

In addition, late HMR remains of prognostic
importance after liver transplantation, with larger area
under the receiver-operating curve than clinical param-
eters and heart rate variability (AUC: 0.79 vs 0.66 and
0.52, respectively) in univariate analysis.22 However,
multivariate analysis revealed that late HMR has no
additive value to a reference model in predicting
outcome (AUC 0.80 vs 0.79, respectively).22

In the AL type population, very little is known
about the consequences of reduced late HMR. Follow-up
of the available studies in this population is too limited
to identify arrhythmogenic consequences of impaired
cardiac sympathetic innervation.8,10,12

Data on the contribution of reduced late HMR to
cardiovascular outcome measurements in patients with
ATTR amyloidosis seems to be incomplete. Only one
study reported the association of reduced late HMR with
the presence of ventricular arrhythmia, and the progres-
sion of conduction disturbances after liver transplantation

Figure 2. Example of a 42 year old female patients with hereditary ATTR amyloidosis (TTR-
Tyr114Cys), without cardiac bone tracer accumulation (A), but impaired cardiac sympathetic
innervation (B 15 minutes p.i., C 4 hours p.i.). Late HMR 1.63.
due to continuous amyloid infiltration. Understanding this apparent oxymoron (i.e.: the cessation of progression of cardiac innervation abnormalities despite continuous amyloid infiltration after liver transplantation) will be a challenge for future investigations. As of yet, the actual incidence of ventricular arrhythmia, sudden cardiac death, or appropriate ICD shocks in amyloidosis patients with impaired cardiac sympathetic innervation is not fully elucidated. Therefore, the question whether amyloidosis patients will benefit from prophylactic ICD remains unanswered.

**RELATION TO OTHER NUCLEAR IMAGING MODALITIES IN AMYLOIDOSIS**

In early studies using [I-123]MIBG, amyloidosis patients underwent additional (rest) myocardial perfusion scintigraphy using thallium-201 ([TI-201]).
Table 1. Main results and patient outcome as reported in studies using Iodine-123 labelled metaiodobenzylguanidine scintigraphy in patients with amyloidosis

<table>
<thead>
<tr>
<th>Study</th>
<th>Author, year of publication</th>
<th>Number of patients</th>
<th>Tracer dose</th>
<th>Collimator type</th>
<th>Time point late HMR</th>
<th>Amyloid typing</th>
<th>Main results</th>
<th>Patient outcome</th>
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<tbody>
<tr>
<td>1</td>
<td>Nakata et al⁴⁶</td>
<td>1 patient</td>
<td>111 MBq (3 mCi) [I-123]-MIBG</td>
<td>N/A</td>
<td>4 hours p.i.</td>
<td>hATTR (TTR Val30Met)</td>
<td>No cardiac tracer accumulation</td>
<td>N/A</td>
</tr>
<tr>
<td>2</td>
<td>Tanaka et al⁵⁵</td>
<td>12 patients</td>
<td>148 MBq (4 mCi) [I-123]-MIBG</td>
<td>LE</td>
<td>3 hours p.i.</td>
<td>hATTR</td>
<td>No cardiac tracer accumulation in 8 of 12</td>
<td>Mean FU 15.5 ± 5.8 months: no lethal arrhythmia, no cardiac death</td>
</tr>
<tr>
<td>3</td>
<td>Delahaye et al⁵⁶</td>
<td>17 patients, 12 healthy controls</td>
<td>300 MBq (8 mCi) [I-123]-MIBG</td>
<td>LE</td>
<td>4 hours p.i.</td>
<td>hATTR</td>
<td>Mean late HMR in patients 1.36 ± 0.26 vs in healthy controls 1.98 ± 0.35 (P &lt; 0.001), no difference in wash-out</td>
<td>N/A</td>
</tr>
<tr>
<td>Study</td>
<td>Author, year of publication</td>
<td>Number of patients</td>
<td>Tracer dose</td>
<td>Collimator type</td>
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<tr>
<td>4</td>
<td>Delahaye et al&lt;sup&gt;35&lt;/sup&gt;</td>
<td>21 patients, 12 healthy controls</td>
<td>150 and 180 MBq (4 and 5 mCi) [C-11]-MQNB and 300 MBq (8 mCi) [I-123]-MIBG</td>
<td>LE</td>
<td>4 hours p.i.</td>
<td>hATTR (20 patients TTR Val30Met, 1 patient TTR Thr49Ala)</td>
<td>Mean muscarinic receptor density was higher in patients than in control subjects: B'max, 35.5 ± 8.9 vs 26.1 ± 6.7 pmol/mL (P = 0.003) Mean late HMR in patients 1.43 ± 0.28 vs in healthy controls 1.98 ± 0.35 (P &lt; 0.001), mean wash-out 29% ± 6.8% vs 21% ± 6% (P = 0.003). Individual muscarinic receptor density did not correlate with late HMR</td>
<td>N/A</td>
</tr>
<tr>
<td>5</td>
<td>Watanabe et al&lt;sup&gt;9&lt;/sup&gt;</td>
<td>4 patients, 10 age-matched controls</td>
<td>111 MBq (3 mCi) [I-123]-MIBG</td>
<td>N/A</td>
<td>4 hours p.i.</td>
<td>hATTR (TTR Val30Met)</td>
<td>Mean late HMR in patients 1.1 ± 0.2, vs 2.4 ± 0.2 in health controls (p-value N/A)</td>
<td>N/A</td>
</tr>
</tbody>
</table>
Table 1 continued

<table>
<thead>
<tr>
<th>Study Author, year of publication</th>
<th>Number of patients</th>
<th>Tracer dose Collimator type</th>
<th>Time point late HMR</th>
<th>Amyloid typing</th>
<th>Main results</th>
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<tbody>
<tr>
<td>6 Hongo et al8</td>
<td>25 patients, of which 16 patients without and 9 patients with autonomic neuropathy</td>
<td>111 MBq (3 mCi) [I-123]-MIBG</td>
<td>LE</td>
<td>3 hours p.i.</td>
<td>AL</td>
<td>Mean late HMR in patients without autonomic neuropathy $1.53 \pm 0.06$ vs in with autonomic neuropathy $1.29 \pm 0.05$ ($P &lt; 0.001$), mean wash-out $42 \pm 4.8%$ vs $31 \pm 3.8%$ ($P &lt; 0.001$)</td>
</tr>
<tr>
<td>7 Lekakis et al10</td>
<td>3 patients, 23 controls</td>
<td>185 MBq (5 mCi) [I-123]-MIBG</td>
<td>LE</td>
<td>4 hours p.i.</td>
<td>AL</td>
<td>Mean late HMR $1.33 \pm 0.1$ vs in 2.13 ± 0.2 healthy controls ($P$ value N/A)</td>
</tr>
<tr>
<td>8 Coutinho et al28</td>
<td>34 patients, of which 2 patients without and 12 patients with autonomic neuropathy</td>
<td>[I-123]-MIBG (dose N/A)</td>
<td>N/A</td>
<td>N/A</td>
<td>hATTR</td>
<td>Mean late HMR $1.75 \pm 0.5$ in all patients. Mean late HMR in patients without neuropathy $2.2 \pm 0.5$ vs patients with neuropathy $1.5 \pm 0.4$ ($P = 0.001$)</td>
</tr>
<tr>
<td>9 Delahaye et al11</td>
<td>31 patients</td>
<td>300 MBq (8 mCi) [I-123]-MIBG</td>
<td>LE</td>
<td>4 hours p.i.</td>
<td>hATTR</td>
<td>Mean late HMR 2 years after liver transplantation $1.46 \pm 0.28$ vs 6 months before liver transplantation $1.45 \pm 0.29$, $P$ = not significant</td>
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</tbody>
</table>

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<tr>
<td>10</td>
<td>Algalarrando et al\textsuperscript{36}</td>
<td>32 patients</td>
<td>300 MBq (8 mCi) \textsuperscript{[I-123]}-MIBG</td>
<td>LE 4 hours p.i.</td>
<td>hATTR</td>
<td>Late HMR $\leq$ 1.6 in 26 out of 32 patients</td>
<td>No cardiac death or lethal arrhythmia reported</td>
</tr>
<tr>
<td>11</td>
<td>Noordzij et al\textsuperscript{12}</td>
<td>61 patients, 9 healthy control subjects</td>
<td>185 MBq (5 mCi) \textsuperscript{[I-123]}-MIBG</td>
<td>ME 4 hours p.i.</td>
<td>AL (39 patients), AA (11 patients), ATTR (11 patients)</td>
<td>Mean late HMR in all patients 2.3 ± 0.75 vs healthy control subjects 2.9 ± 0.58 ($P &lt; 0.005$). Mean late HMR in ATTR patients 1.7 ± 0.75 vs AL patients 2.4 ± 0.75 ($P &lt; 0.05$). Mean wash-out in patients 8.6% ± 14% vs in healthy control subjects 2.1% ± 10% ($P &lt; 0.05$)</td>
<td>No cardiac death or lethal arrhythmia</td>
</tr>
<tr>
<td>12</td>
<td>Noordzij et al\textsuperscript{37}</td>
<td>2 patients</td>
<td>185 MBq (5 mCi) \textsuperscript{[I-123]}-MIBG</td>
<td>ME 4 hours p.i.</td>
<td>wtATTR, hATTR (TTR Val122Ile)</td>
<td>Patient A: late HMR 1.57, wash-out &gt;20%, patient B: late HMR 1.13, wash-out 28%</td>
<td>N/A</td>
</tr>
<tr>
<td>13</td>
<td>Coutinho et al\textsuperscript{21}</td>
<td>143 patients</td>
<td>185 MBq (5 mCi) \textsuperscript{[I-123]}-MIBG</td>
<td>LE 3 hours p.i.</td>
<td>hATTR (TTR Val30Met)</td>
<td>Mean late HMR 1.83±0.43, and mean was-out 47±11%</td>
<td>Mean FU 5.5 years: hazard ratio all-cause mortality 7 if HMR &lt;1.6, progressive increase in 5-year mortality with decrease in late HMR</td>
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<td>Study</td>
<td>Author, year of publication</td>
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<td>Tracer dose</td>
<td>Collimator type</td>
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<tr>
<td>14</td>
<td>Takahashi et al[^3^]</td>
<td>6 patients</td>
<td>[I-123]-MIBG (dose N/A)</td>
<td>N/A</td>
<td>N/A</td>
<td>hATTR (TTR Val30Met)</td>
<td>Mean late HMR at baseline 1.7 ± 0.9 vs after 3 years diflunisal treatment 1.9 ± 1.0 (P = 0.004). Mean wash-out at baseline 46% ± 20% vs after 3 years 43% ± 23% (P = 0.67)</td>
</tr>
<tr>
<td>15</td>
<td>Algalarrando et al[^2^]</td>
<td>215 patients</td>
<td>3 MBq/kg (0.08 mCi/kg) [I-123]-MIBG</td>
<td>LE</td>
<td>4 hours p.i.</td>
<td>hATTR (148 patients TTR Val30Met)</td>
<td>Median late HMR 1.49 (Inter-quartile range 1.24 - 1.74, range 0.97 - 2.52)</td>
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<tr>
<td>Study</td>
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<td>Tracer dose</td>
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<tr>
<td>16</td>
<td>Azevedo Coutinho et al&lt;sup&gt;23&lt;/sup&gt;</td>
<td>232 patients</td>
<td>185 MBq (5 mCi) [I-123]-MIBG</td>
<td>LE</td>
<td>3 hours p.i.</td>
<td>hATTR (TTR Val30Met)</td>
<td>Initial assessment: mean late HMR 1.83 ± 0.03, median wash-out 2.5 (Inter-quartile range −2.3–8.5)</td>
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</table>

[I-123]-MIBG, Iodine-123 labelled metaiodobenzylguanidine; [C-11]-MQNB, carbon-11 labelled methylquinuclidinyl benzilate; Hattr, hereditary transthyretin-derived amyloid; wtATTR, wild-type transthyretin-derived amyloid; AL, immunoglobulin light chain-derived amyloid; FU, follow-up; HMR, heart-to-mediastinum ratio; HR, hazard ratio; LE, low energy; ME, medium energy; N/A, not available; p.i., post injection
None of the included patients seemed to suffer from myocardial infarction, since all rest [Tl-201] scans were reported normal, without perfusion defects. This perfusion – innervation mismatch is a known phenomenon in patients with ischemic cardiomyopathy, but also occurs in patients with non-ischemic (dilating) cardiomyopathy.\textsuperscript{24,25} Myocardial perfusion abnormalities are known to result in damaged sympathetic nerve terminals, leading to a larger area of impaired innervation than impaired perfusion alone. This mismatch pattern leads to electrophysiological imbalance, which is associated with a higher risk of developing ventricular dysrhythmia.\textsuperscript{24,25} The mechanism behind the development of perfusion–innervation mismatch pattern in patients with non-ischemic cardiomyopathy is not fully elucidated. However, the presence of structural changes (for example heterogeneous interstitial fibrosis) may contribute to altered ventricular activation and contractility, due to maladaptation to myocardial injury. In combination with disturbed sympathetic stimulation due to amyloid infiltration, this may contribute to a higher risk of ventricular dysrhythmia in amyloidosis patients as well.

The mutual contribution of autonomic neuropathy and cardiomyopathy to each other on decreased late HMR remains a conundrum. Since both wtATTR and hATTR type amyloidosis patients show decreased late HMR, \textsuperscript{[I-123]MIBG} scintigraphy alone may not be sufficient to discriminate between autonomic neuropathy and cardiomyopathy. Several studies have shown that myocardial bone tracer accumulation discriminates hATTR from AL type amyloidosis.\textsuperscript{26,27} Bone tracer accumulation predominantly occurs in wild-type ATTR type patients, probably as a result of the underlying cardiomyopathy. On the contrary, patients with hATTR type amyloidosis without cardiomyopathy tend to show no myocardial bone tracer accumulation, and normal biomarkers (N-terminus pro-brain natriuretic peptide, and troponine-T). Within these patients, late HMR is generally lower in the subgroup of patients with other symptoms of polyneuropathy.\textsuperscript{26} Future studies should focus on the possible additive value of bone scintigraphy in relation to \textsuperscript{[I-123]MIBG} scintigraphy in getting a better understanding of the mutual contribution of neuropathy and cardiomyopathy to each other in ATTR type patients.

Recently in positron emission tomography (PET), carbon-11 labelled Pittsburgh compound-B ([C-11]-PiB), derived from the amyloid stain thioflavin, as well as fluorine-18 ([F-18]) labelled florbetapir have been used as tracers for cardiac amyloid.\textsuperscript{29,30} However, their role against cardiac sympathetic innervation is to be determined. There is no role for [F-18] fluorodeoxyglucose (FDG) imaging or \textsuperscript{[I-123]SAP} scintigraphy in evaluating cardiac manifestation against sympathetic innervation disturbances in amyloidosis, since neither one of both tracers is known to accumulate in cardiac amyloid deposits.\textsuperscript{31,32}

### FUTURE DEVELOPMENTS

There is an increasing evidence for the prognostic value of \textsuperscript{[I-123]MIBG} scintigraphy in patients with amyloidosis. However, more prospectively acquired data is needed to implement \textsuperscript{[I-123]MIBG} scintigraphy in guidelines as a standard imaging procedure in the management of (especially ATTR type) amyloidosis patients. Therefore, consensus in acquisition parameters in different study protocols is pivotal. Standardization of collimator choice, imaging acquisition, and data analysis in different studies, is necessary for successful implementation in daily patient practice.\textsuperscript{15}

Finally, the use of PET tracers has advantages over \textsuperscript{[I-123]MIBG} in cardiac sympathetic innervation imaging. Carbon-11 labelled meta-hydroxy-ephedrine [C-11]mHED has been extensively studied in patients with both ischemic and non-ischemic cardiomyopathies.\textsuperscript{14,20} Based on the studies in patients with left ventricular dysfunction, [C-11]mHED outperforms \textsuperscript{[I-123]MIBG} in detecting regional impaired sympathetic innervation, due to better resolution and absolute quantification.\textsuperscript{33} Despite that [C-11]mHED is the most used PET tracer for visualization of cardiac sympathetic innervation abnormalities, it’s value has not yet been studied in amyloidosis patients. Future studies should provide information on the value of recently developed PET tracers in evaluating cardiac sympathetic innervation in amyloidosis patients. In theory, two new PET tracers may have additional value over \textsuperscript{[I-123]MIBG} scintigraphy in regard to higher HMR. For example, \textsuperscript{[I-124]MIBG} may provide superior image quality, whereas N-[3-Bromo-4-3-[F-18]fluoro-propoxy]-benzyl]-guanidine ([F-18]LM1195) has the additional advantage that an on-site cyclotron is not necessary.\textsuperscript{34}

### CONCLUSIONS

\textsuperscript{[I-123]MIBG} is currently the most widely used radiopharmaceutical for imaging cardiac sympathetic innervation disturbances in patients with cardiac manifestations of amyloidosis. Particular patients with hATTR type amyloidosis show diminished late HMR’s, and consequently have a higher risk of cardiac mortality.

Future studies should provide better insight into the presence and degree of overlap between cardiac neuropathy and cardiomyopathy in patients with cardiac manifestations of amyloidosis, the role of nuclear medicine modalities in distinguishing cardiac neuropathy from cardiomyopathy, and finally, the potential role
of PET tracers in evaluating impaired cardiac sympathetic innervation.

Disclosure

All Authors have no disclosure to state.

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