Myocardial Emission Computed Tomography with Iodine-123-Labeled Beta-Methyl-Branch Fatty Acid in Patients with Hypertrophic Cardiomyopathy

Chinori Kurata, Kei Tawarahara, Takahisa Taguchi, Shigeyuki Aoshima, Akira Kobayashi, Noboru Yamazaki, Hiroaki Kawai, and Masao Kaneko

Third Department of Internal Medicine and Department of Radiology, Hamamatsu University School of Medicine, Hamamatsu, Japan

We studied whether emission computed tomography (ECT) with 123I-labeled 15-(p-iodophenyl)-3R,S-methylpentadecanoic acid (BMIPP) shows myocardial distribution different from 201TI in hypertrophic cardiomyopathy. In 10 patients with asymmetric septal hypertrophy (ASH), 5 with diffuse hypertrophy and 2 with apical hypertrophy, ECT was acquired 30 min (early) and 4 hr (late) after injection of 123I-BMIPP at rest and was compared with ECT with 201TI. In 10 patients with ASH, the relative regional uptake (RRU) of the septum was lower in the early 123I-BMIPP study than in the 201TI study, although that of the posterior wall was similar. In the early and late 123I-BMIPP studies, the RRU of the septum was lower in 10 patients with ASH than in 7 without ASH, although that of the posterior wall was similar. In the 201TI study, however, the RRU of both the septum and posterior wall was similar in those with and without ASH. Moreover, in 16 patients, the apparent left ventricular size was larger in the early 123I-BMIPP study than in the 201TI study, which suggested reduced 123I-BMIPP uptake in the subendocardium. In patients with hypertrophic cardiomyopathy, thus, 123I-BMIPP imaging may reveal impaired regional fatty acid utilization, which is independent of regional perfusion.


Myocardial metabolic imaging has been expected not only to enhance our understanding of cardiomyopathies, but also to aid in the development of effective treatment of cardiomyopathies (1). Recently, several studies using PET have demonstrated regional changes in myocardial metabolism in patients with cardiomyopathy (2-6). Unfortunately, positron imaging is not widely available at present. Several radioiodinated fatty acids have been used for investigating myocardial accumulation and turnover of fatty acids in patients with dilated cardiomyopathy, ischemic heart disease and hypertensive hypertrophy (7-12). It, however, remains unclear whether metabolic imaging with radioiodinated fatty acids can elucidate the characteristic features of patients with hypertrophic cardiomyopathy (HCM).

Myocardial perfusion imaging with 201TI has demonstrated regional perfusion abnormalities in patients with various cardiomyopathies (13-18). Perfusion defects in those patients might represent myocardial scar and/or ischemia. If an uncoupling of regional perfusion and fatty acid utilization exists in the myocardium of cardiomyopathy, myocardial imaging with radioiodinated fatty acids could elucidate unique features different from 201TI imaging. In the cardiomyopathic Bio 14.6 Syrian hamster, an animal model for HCM, we have demonstrated discrepancies in the myocardial distributions of 201TI and 123I-labeled 15-(p-iodophenyl)3R,S-methylpentadecanoic acid (BMIPP) (19). The present study was therefore undertaken to investigate whether emission computed tomography (ECT) with 123I-BMIPP may show myocardial distribution different from that with 201TI in patients with HCM.

METHODS

Patients Characteristics

The study group consisted of 17 patients (11 men and 6 women), ranging in age from 21 to 75 yr (Table 1). In all 17 patients, left ventricular hypertrophy was diagnosed on echocardiographic demonstration. Two of these 17 patients had coronary artery disease; one had a significant stenosis of the right coronary artery, hypertension and diabetes mellitus; and the other patient had three-vessel disease with previous myocardial infarction. The two patients, however, were diagnosed as having HCM as a complication because their hypertrophy, limited to the interventricular septum, could not be explained by coronary artery disease alone. The remaining 15 of these 17 patients had normal coronary angiograms and were diagnosed as having HCM because their left ventricular hypertrophy could not be ascribed to another cardiac or systemic disease.
SPECT with Fatty Acid Analogue in HCM • Kurata et al

Echocardiographic Studies

Echocardiographic studies were performed within a week of $^{123}$I-BMIPP imaging, using a phased-array scanner operating at 2.5 or 3.5 MHz (Model SSD870, Aloka, Tokyo, Japan). The M-mode echocardiographic examination was performed by moving an M-mode cursor on the two-dimensional parasternal long-axis view. The M-mode echocardiographic measurements were performed according to the criteria recommended by the American Society of Echocardiography (20) and the results are shown in Table 1.

Asymmetric septal hypertrophy (ASH) was considered to be present if the end-diastolic thickness of the septum was at least 15 mm and its ratio to that of the left ventricular posterior wall was at least 1.3. Diffuse hypertrophy was considered to be present if both of the end-diastolic thicknesses of the septum and posterior wall were at least 15 mm. If echocardiographic apical long-axis or four-chamber view demonstrated apical hypertrophy and a characteristic spade-like configuration was demonstrated in the right anterior oblique ventriculogram at end-diastole (21), apical hypertrophy was considered to be present. According to these criteria, 10 had ASH, 5 had diffuse hypertrophy and 2 had apical hypertrophy (Table 1). The two patients with coronary artery disease (Patients 9 and 10) had ASH.

ECT Acquisitions

All patients gave informed consent to a protocol of $^{123}$I-BMIPP imaging as a Phase 2 clinical trial. The study protocol and consent form had been approved by the Hamamatsu University School of Medicine Human Subject Protection Committee. On the day of $^{123}$I-BMIPP imaging, patients had no breakfast and continued fasting until the end of the late imaging session. At rest, 111 MBq (3 mCi) of $^{123}$I-BMIPP were injected intravenously. Early tomographic imaging was started 30 min after the injection, using a large field of view rotating gamma camera equipped with a low-energy, all-purpose collimator centered on the 159 keV photo peak with a 20% window. The camera was rotated over 180 degrees from the 45-degree right anterior oblique to the 45-degree left posterior oblique position. Thirty-two images were obtained in a 64 × 64 matrix for 30 sec per image on a dedicated nuclear medicine computer (Scintipac 2400, Shimadzu, Kyoto, Japan). No attenuation or scatter correction was used. Late imaging was performed 4 hr after injection using the same imaging protocol as that for early imaging.

Four or six days after the $^{123}$I-BMIPP studies, $^{201}$Tl imaging was performed after an overnight fast. At rest, 111 MBq (3 mCi) of $^{201}$Tl were injected intravenously. Emission computed tomograms were acquired 10 min after injection using the same imaging protocol as that for early imaging. No attenuation or scatter correction was used. Late imaging was performed after an overnight fast.

For each of the $^{201}$Tl and early and late $^{123}$I-BMIPP acquisition data, the same reconstruction of tomographic images was performed. Each data set was reconstructed by a Shepp-Logan filtered backprojection algorithm after preprocessing of the projection images with Butterworth filter correction as 1-pixel thick transverse slices. Vertical long-axis, short-axis and horizontal long-axis slices, each 6 mm thick, were reconstructed from the transverse slices and were magnified two diameters using the same left ventricular oblique angles for each of the $^{201}$Tl and early and late $^{123}$I-BMIPP data.

Quantitative Analysis of Tomograms

From each set of the $^{201}$Tl and early and late $^{123}$I-BMIPP short-axis slices, three composite slices 18 mm or 24 mm thick were constructed by adding three or four short-axis slices, respectively. The same number of short-axis slices (three or four) was used to reconstruct the composite slices for each of the $^{201}$Tl and early and late $^{123}$I-BMIPP studies in each patient. The three composite slices consisted of basal, mid- and apical ventricular levels. The composite basal and mid-ventricular slices were divided, respectively, into eight segments, and the composite apical ventricular slice was divided into five segments (Fig. 1). A square region of interest of 5 × 5 pixels was placed over the center of each of the

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HCM = hypertrophic cardiomyopathy; IVSth = end-diastolic thickness of the septum (mm); LVPWth = end-diastolic thickness of the posterior wall (mm); LVDD = end-diastolic left ventricular dimension (mm); LVDs = end-systolic left ventricular dimension (mm); LVEF = left ventricular ejection fraction (%); and ASH = asymmetric septal hypertrophy.
total 21 segments. For each segment, the regional uptake was determined for each segment as mean counts per pixel within the corresponding region of interest. Moreover, the relative regional uptake (RRU, %) was determined for each segment as the ratio of its regional uptake to the maximum regional uptake within the total 21 segments of each set of the three composite short-axis slices. In the $^{123}$I-BMIPP studies, the washout rate (%) in each segment was calculated by subtracting regional uptake in the late image from the corresponding regional uptake in the early image and dividing by that in the early image.

**Apparent Left Ventricular Size**

In our previous experimental study using cardiomyopathic hamsters, more $^{123}$I-BMIPP accumulated in the epicardial side of the left ventricular myocardium than in the endocardial side, although $^{201}$Tl accumulated almost equally in the epicardial and endocardial sides (19). If the experimental findings could be applied to human cardiomyopathy, the apparent size of the left ventricular cavity on the $^{123}$I-BMIPP tomogram might be larger than that on the $^{201}$Tl tomogram. The apparent left ventricular size was therefore compared between the $^{201}$Tl and early $^{123}$I-BMIPP tomograms, using the following method. A short-axis slice in the mid- to basal ventricular level from each of the $^{201}$Tl and early $^{123}$I-BMIPP studies was chosen so that two short-axis slices from the two studies anatomically corresponded to each other. In the short-axis slice, the center was assigned from which 60 equidistant radii (6 degrees apart) were generated toward the left ventricular periphery. A region of interest was generated by connecting 60 points with the maximum counts per pixel within each radius. The apparent left ventricular size was evaluated with the number of pixels within the region of interest.

**Statistical Methods**

The data are presented as mean ± s.d. The statistical significance of differences in mean values between two groups was analyzed with the paired or unpaired Student’s t-test. When test groups were not normally distributed, the paired or unpaired nonparametric Wilcoxon t-test was applied. Correlations between two variables were examined using linear regression analysis. Probability (p) values of <0.05 (two-sided) were considered to be statistically significant.

**RESULTS**

**Regional Uptakes of $^{201}$Tl and $^{123}$I-BMIPP**

In all of the early and late studies, $^{123}$I-BMIPP provided high contrast tomograms of the left ventricular myocardium (Fig. 2), although the quality of the late $^{123}$I-BMIPP images was relatively poor compared with the $^{201}$Tl and early $^{123}$I-BMIPP images.

The mean regional uptake of the total 357 segments (17 patients × 21 segments/patient) was 445 ± 216, 228 ± 81 and 133 ± 38 counts/pixel in the $^{201}$Tl and early and late $^{123}$I-BMIPP studies, respectively. The regional uptakes of $^{201}$Tl correlated with those of $^{123}$I-BMIPP in the early study ($r = 0.781; p < 0.001$) and in the late study ($r = 0.664; p < 0.001$). The regional uptakes of $^{123}$I-BMIPP in the late study correlated with those in the early study ($r = 0.904; p < 0.001$). Regional uptake was significantly higher in the $^{201}$Tl study than in the early $^{123}$I-BMIPP study ($p < 0.001$).

In the 357 segments, the washout rate of $^{123}$I-BMIPP from the early to the late study was 40.1% ± 9.4%.

**Relative Regional Uptake of $^{201}$Tl and $^{123}$I-BMIPP**

In the 357 segments, the RRU was 75.5% ± 15.4%, 75.8% ± 15.2% and 77.5% ± 14.3% in the $^{201}$Tl and early and late $^{123}$I-BMIPP studies, respectively. The RRU of $^{201}$Tl correlated with those of $^{123}$I-BMIPP in the early study ($r = 0.714; p < 0.001$) and in the late study ($r = 0.618; p < 0.001$). The RRU of $^{123}$I-BMIPP in the late study correlated with that in the early study ($r = 0.863; p < 0.001$).

The mean difference in the RRU between the $^{201}$Tl and early $^{123}$I-BMIPP studies was 0.3% ± 11.6% (RRU of $^{123}$I-BMIPP minus RRU of $^{201}$Tl). The distribution of the differences in the 357 segments is shown in Figure 3A. The absolute value of the difference in the RRU between the $^{201}$Tl and early $^{123}$I-BMIPP studies was greater than 10% in 146 (41%) of the 357 segments and than 20% in 29 segments (8%). Similarly, the mean difference in the
A RELATIVE REGIONAL UPTAKE (Early-TO-10~5-lOll, 1~5~21y/l,
ZIRELATIVE REGIONAL UPTAKE (Late- Early)
100.

FIGURE 3. (A) Distribution of the difference in relative regional
uptake (RRU) between the 2°1TI and early 123I-BMIPP studies in
357 segments. The RRU of 123I-BMIPP in a segment minus that
of 2°1TI in the corresponding segment (∆ Relative Regional Uptake
(Early- TII)) on the x-axis is classified into six levels: -< -30%
(-40<, =<-30%), -< -20% (-30%<, -< -20%), -10% ~ 10% (-10%<, <10%), 10%~< (10%~<, <20%),
and 20%~< (20%~<, <30%). Hatched bars represent segments
with the absolute value of the ∆ relative regional uptake greater
than 10%. (B) Distribution of the difference in RRU between the
early and late 123I-BMIPP studies. The RRU in the late study
minus that in the early study (A Relative Regional Uptake (Late-
Early)) is classified in the same manner as in A above.

RRU between the early and late 123I-BMIPP studies was
1.8% ± 7.7% (RRU in the late study minus RRU in the
early study). The distribution of the differences in the 357
segments is shown in Figure 3B. The absolute value of the
difference in the RRU between the early and late 123I-
BMIPP studies was greater than 10% in 69 (19%) of the
357 segments and than 20% in 6 segments (1.7%).

Apparent Size of the Left Ventricle
The left ventricular cavity was apparently larger in the
123I-BMIPP study than in the 2°1TI study, particularly in
patients with diffuse hypertrophy (Fig. 4). The ratio of the
apparent left ventricular size in the early 123I-BMIPP study
to that in the 2°1TI study was 1.20 ± 0.03 in 17 patients.
The ratio in the 16 patients was greater than 1.0. To
examine whether the finding might be due to an artifact
of reconstruction or attenuation, we performed an
experiment using a left ventricular myocardium phantom and
the same imaging protocol as in the clinical study. As a
result, the ratio of apparent ventricular sizes in the 123I-
BMIPP study to that in the 2°1TI study was 1.03 ± 0.03,
which was significantly smaller than that in the 17 patients
(p < 0.05). Moreover, in 14 patients (82%), the ratio was
greater than 1.09 (mean ± 2 s.d. in the phantom experiment).

Iodine-123-BMIPP Tomograms in Patients with ASH
In the 10 patients with ASH, the RRU of the septum
(the mean of Segments 3 and 11) was 76% ± 8%, 70% ±
9% and 70% ± 11%, and the RRU of the posterior wall
(the mean of Segments 6 and 14) was 75% ± 15%, 76%
± 15% and 78% ± 13% in the 2°1TI and early and late 123I-
BMIPP studies, respectively. The RRU of the septum in
the early 123I-BMIPP study was lower than that in the 2°1TI
study (p < 0.05), although that of the posterior wall was
similar in the two studies (p > 0.05). However, the RRU
of both the septum and posterior wall was similar in the
2°1TI and late 123I-BMIPP studies (p > 0.05, respectively).
Figure 5 shows examples of the patients with ASH.

In the seven patients without ASH, the RRU of the
septum was 79% ± 11%, 86% ± 7% and 86% ± 8%, and
the RRU of the posterior wall was 76% ± 5%, 77% ± 4%
and 80% ± 6% in the 2°1TI and early and late 123I-BMIPP
studies, respectively. Figure 6 shows the comparison of
RRU between the 10 patients with and the 7 without ASH.
In both the early and late 123I-BMIPP studies, the RRU
of the septum was lower in the 10 patients with ASH than
in the 7 without ASH (p < 0.01 and p < 0.05, respectively),
although that of the septum in the 2°1TI study was similar
(p > 0.05, respectively). In all of the 2°1TI and early and
late 123I-BMIPP studies, however, the RRU of the posterior
wall was similar in those with and without ASH (p > 0.05,
respectively).

Furthermore, the ratio of the RRU in the early 123I-
BMIPP study to that in the 2°1TI study was compared
between the 10 patients with and the 7 without ASH (Fig.
7). The ratio in the septum was smaller in the 10 patients
with ASH (0.92 ± 0.09) than in the 7 without ASH (1.09
± 0.10; p < 0.01). However, the ratio of the posterior wall

FIGURE 4. Tomograms of Patient 12 with diffuse hypertrophy.
The left ventricle appears to be larger in the early 123I-BMIPP
images than in the 2°1TI images. The difference suggests that
transmural distribution may not be similar in the two tracers.
These tomograms also shows a matched decrease of 2°1TI and
123I-BMIPP in the subendocardium of the inferolateral wall. How-
ever, it is not certain because the spatial resolution of this imaging
is too poor to discriminate the endocardium from the epicardium.
Abbreviations as in Figure 2.
FIGURE 5. Tomograms of Patients 5A and 6B with septal hypertrophy. (A) The septal uptake (arrows) is high in the 20T1 images but markedly low in the early 123I-BMIPP images. (B) The uptake of 123I-BMIPP (arrows) is lower in the septum than in the other regions, although that of 20T1 is not so different. Abbreviations as in Figure 2.

was similar in those with and without ASH (1.03 ± 0.10 versus 1.02 ± 0.07; p > 0.05).

Thus, in the patients with ASH, the disproportionately thickened septum showed reduced uptake of 123I-BMIPP compared to 20T1. In the two patients with ASH and congestive heart failure, however, the 20T1 and 123I-BMIPP tomograms demonstrated similar defects (Fig. 8).

Iodine-123-BMIPP Tomograms in Patients with Diffuse or Apical Hypertrophy

The five patients with diffuse hypertrophy showed relatively similar myocardial distributions of activity in the 20T1 and early and late 123I-BMIPP tomograms, except for the apparent difference in left ventricular size (Figs. 2 and 4). Of the two patients with apical hypertrophy, one showed a 123I-BMIPP defect in the apical region where the 20T1 uptake was higher compared with the other regions (Fig. 9), although the other did not show such a discrepancy between 20T1 and 123I-BMIPP uptake in the apical region.

DISCUSSION

We have demonstrated that patients with HCM showed a discrepancy of myocardial distributions between 20T1 and 123I-BMIPP.

Iodine-123-BMIPP Tomograms in Patients with ASH

The disproportionately thickened septum in the patients with ASH showed a reduced uptake of 123I-BMIPP independent of regional perfusion assessed by 20T1, although such an uncoupling of 20T1 and 123I-BMIPP was not detected in those without ASH. Moreover, the patients...
Mechanisms of Myocardial $^{123}$I-BMIPP Uptake

BMIPP was designed to have prolonged myocardial retention suitable for ECT (27). The insertion of a methyl radical in the beta-position inhibits beta oxidation and prolongs myocardial retention. The myocardial uptake of BMIPP, therefore, does not directly reflect beta-oxidation of fatty acids in the myocardium. In fact, recent experimental studies using carnitine acyltransferase I inhibitors, 2-tetradecylglycidic acid (28) or 2[5(4-chlorophenyl)pentyloxirane-2-carboxylate (29), demonstrated that the myocardial extraction of BMIPP was not decreased by these inhibitors.

Myocardial accumulation of BMIPP has been reported to be observed mainly in the triglyceride fraction (30) and to be associated with the synthesis of triglyceride (28). Furthermore, it has been demonstrated that myocardial BMIPP accumulation may be closely correlated with intracellular concentrations of adenosine triphosphate, which is required in the first step of enzymatic conversion of fatty acids to acyl-CoA, a common pathway of fatty acid metabolism, such as triglyceride synthesis and $\beta$-oxidation (31). Thus, the uptake of BMIPP may reflect some aspect of myocardial fatty acid metabolism, such as incorporation into triglyceride storage products or an oxidative process responsible for the conversion of BMIPP to polar catabolites (32).

Mechanisms of Abnormal $^{123}$I-BMIPP Distribution in HCM

Our finding indicates that the disproportionately thickened septum may have some regional impairment in fatty acid metabolism independent of perfusion. Grover-McKay et al. (6), using PET, reported that $^{11}$C-palmitate uptake was reduced in the disproportionately thickened septum of patients with HCM. They, however, suggested that the septal fatty acid metabolism was normal since the reduced $^{11}$C-palmitate in the septum was most likely a function of decreased blood flow, and the clearance halftime of the early rapid phase and the residual fraction of $^{11}$C-palmitate were similar in the septum and the lateral wall. The difference in septal fatty acid uptake between our study and theirs may be in part due to the difference between the metabolic tracers: $^{11}$C-palmitate can be metabolized via beta-oxidation, while $^{123}$I-BMIPP hardly can be metabolized.

Grover-McKay et al. (6), demonstrated moreover that $^{18}$F-2-deoxyglucose uptake was lower in the septum than in the lateral wall. The decrease in septal glucose utilization accompanied by the flow-dependent decrease in $\beta$-oxidation of fatty acids may result in a decreased myocardial adenosine triphosphate concentration, which may partly explain the reduced septal uptake of $^{123}$I-BMIPP (31,33).

Miller et al. (10) reported that the severely ischemic myocardium demonstrated an excess of BMIPP accumulation over perfusion measured with $^{201}$TI. They suggested that it was due to abnormal retention of BMIPP. The lesser accumulation of $^{123}$I-BMIPP than $^{201}$TI observed in our

Transmural Differences of $^{123}$I-BMIPP Uptake

The apparent left ventricular size was larger in the ECT with $^{123}$I-BMIPP compared to $^{201}$TI. It suggests that the hypertrophied myocardium may accumulate less $^{123}$I-BMIPP in the subendocardium than in the subepicardium. Using the count peaks of profile curves on ECT with $^{201}$TI, Takeishi et al. (22) measured an apparent left ventricular size and compared it with left ventricular volume measured by radionuclide angiography. They concluded that dipyridamole-induced dilatation of the left ventricular cavity reflected relative subendocardial hypoperfusion. However, it has not been confirmed whether the difference in size between ECT with different isotopes such as $^{201}$TI and $^{123}$I may reflect their different distributions. Comparison between ECT with $^{201}$TI and $^{123}$I-BMIPP in normal controls is necessary for concluding that their transmural distributions are different than that in patients with HCM.

Comparison with Autoradiographic Studies

In previous autoradiographic studies using cardiomyopathic hamsters (19,23) or hypertensive rats (24-26), the uptake of methyl-branched fatty acids was lower in the subendocardial region than in the subepicardial region, although perfusion was similar in the two regions. In patients with HCM, similarly, the apparently larger left ventricle in the $^{123}$I-BMIPP study may suggest reduced uptake of $^{123}$I-BMIPP in the subendocardial region without transmural differences in perfusion.
patients is likely to be neither due to flow-dependent reduction in \(^{123}\)I-BMIPP delivery nor to abnormal retention of \(^{123}\)I-BMIPP induced by severe ischemia. The uncoupling of \(^{201}\)TI and \(^{123}\)I-BMIPP in HCM may not reflect secondary changes induced by ischemia but the intrinsic impairment of myocardial metabolism.

**Limitations of the Study**

Our study has several limitations. First, \(^{123}\)I-BMIPP imaging was not performed in healthy subjects since our study was performed as a Phase 2 clinical trial. We thus could not compare the \(^{123}\)I-BMIPP images in patients with HCM with those in normal controls. Earlier experimental studies using \(\beta\)-methyl fatty acid (19,24–26), however, showed that its myocardial distribution was almost consistent with that of regional perfusion in normal animals. Furthermore, the Phase 1 trial of myocardial imaging with \(^{123}\)I-BMIPP in Japan revealed that the myocardial distribution of \(^{123}\)I-BMIPP in normal volunteers was as homogeneous as that of \(^{201}\)TI (34).

Second, we could not evaluate absolute myocardial uptakes of \(^{123}\)I-BMIPP, due mainly to conventional ECT. If they had been measured, myocardial \(^{123}\)I-BMIPP imaging in patients with HCM might have shown not only regional heterogeneity but also a global abnormality of uptake.

Third, our patients did not form a homogeneous group but consisted of three types of HCM. They ranged in age from 21 to 75 yr. Patient 9, an elderly patient with a history of severe hypertension, might represent hypertensive HCM of the elderly (35,36). Further studies are needed to clarify the characteristic features of individual types of HCM (37).

Fourth, strict comparison between \(^{201}\)TI and \(^{123}\)I-BMIPP images was impossible because they were acquired separately. Simultaneous dual-energy acquisition would provide more precise anatomical comparison. It may, however, cause errors in quantification of the uptake of each tracer (38).

Lastly, we should mention two technical problems. One was a partial volume effect, which is common to all imaging techniques. It causes an underestimation of regional uptakes when myocardial wall thickness is less than twice the spatial resolution of the imaging devices (39). However, in the comparison of RRU between \(^{201}\)TI and \(^{123}\)I-BMIPP for the same region, the partial volume effect cancels out because it is the same for both tracers. The differences observed in our study were therefore unlikely to be artifacts of the partial volume effect. Another was the spatial resolution of our imaging system. The resolution (FWHM = 21 mm) was too poor to discriminate the endocardium from the epicardium. Therefore, we measured the apparent ventricular size using a region of interest generated by the peak points on each radius (22).

**Clinical Implications**

We demonstrated that myocardial ECT with \(^{123}\)I-BMIPP may elucidate characteristic features different from \(^{201}\)TI imaging in patients with HCM. Our observations suggest that ECT with \(^{123}\)I-BMIPP may detect regional abnormalities in patients with HCM earlier than ECT with \(^{201}\)TI. In addition, a combination of \(^{201}\)TI and \(^{123}\)I-BMIPP imaging may be useful for the evaluation of various cardiomyopathies, including ischemic (40), hypertensive and dilated cardiomyopathies.

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Chinori Kurata, Kei Tawarahara, Takahisa Taguchi, Shigeyuki Aoshima, Akira Kobayashi, Noboru Yamazaki, Hiroaki Kawai and Masao Kaneko


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