Meta-Iodobenzylguanidine Imaging: From Standardization to Mortality Risk Models in Heart Failure

Kenichi Nakajima, MD, PhD

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Abstract

\(^{123}\)I-meta-iodobenzylguanidine (MIBG) is a potent prognostic marker of chronic heart failure (CHF). However, inter-institutional variations due to methodological variations required minimization before \(^{123}\)I-MIBG findings could be universally applied to the diagnosis, treatment and prognosis of CHF. Therefore, protocols including data acquisition, setting regions of interest for calculating heart-to-mediastinum ratios (HMR) and cross-calibration of HMR among institutions required standardization. A cross-calibration phantom was introduced to overcome institutional differences, and a large amount of experimental data were collected, which enabled multicenter comparisons and the creation of large-scale prognostic databases. Thereafter, cardiac mortality risk models to estimate short- and long-term (two and five years, respectively) mortality were created based on a standardized \(^{123}\)I-MIBG HMR. The ability of these models to accurately determine prognosis is currently undergoing validation.

Keywords: Calibration phantom, Cardiac imaging, Data processing, Heart-to-mediastinum ratio, Prognosis

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Kenichi Nakajima
Department of Nuclear Medicine, Kanazawa University Hospital, 13-1 Takara-machi, Kanazawa, Japan 920-8641
E-mail: nakajima@med.kanazawa-u.ac.jp

The clinical application of \(^{123}\)I-meta-iodobenzylguanidine (MIBG) was approved in Japan during 1992 (1), and it has since become established in the guidelines of the Japanese Circulation Society as an effective means of evaluating the severity, therapeutic effects and prognosis of chronic heart failure (CHF) (2). Quantitation of uptake has supported the effectiveness of \(^{123}\)I-MIBG in clinical practice and in research studies. The heart-to-mediastinum ratio (HMR) is a simple method in which regions of interest (ROIs) are placed on the heart and mediastinum, and then their average ratio is calculated (3). However, simplicity does not necessarily mean reliability, reproducibility or practicality among hospitals, where preferences for data acquisition and processing methods vary considerably. The varying factors among institutions that affect the stability of results include radiotracer doses administered, image acquisition protocols, ROI settings for processing (4, 5), and corrections for camera-collimator differences. Cardiac mortality risk models were also created (6), and these are addressed elsewhere in this article. However, a fluctuating HMR influenced the final prediction of cardiac mortality, which could seriously impact decisions about patient management. Whereas minor differences in institutional preferences might be acceptable, diagnostic instability, differences in assessments of therapeutic effects and prognosis should be minimized before \(^{123}\)I-MIBG could be universally applied. This article therefore addresses which issues involved in \(^{123}\)I-MIBG imaging require standardization.

Administration of \(^{123}\)I-MIBG and data acquisition

The amount of \(^{123}\)I-MIBG (MyoMIBG; FUJIFILM RI Pharma Co. Ltd., Tokyo, Japan) administered for clinical studies in Japan is 111 MBq, which is lower than the amounts of AdreView (GE Healthcare, Little Chalfont, UK; Arlington Heights, IL, USA) applied in the USA (370 MBq) and Europe (185 MBq). Scintigraphic images were acquired as 256 × 256 matrices over 3-10 minutes in the anterior planar view. Early and late images were acquired at 15-30 minutes and 3-4 hours, respectively, after tracer administration, and energy was centered at 159 keV with a 15% or 20% window. One European proposal recommends medium-energy (ME) (4) or...
ME general-purpose (MEGP) collimators, but low-energy high-resolution (LEHR), low-energy general-purpose (LEGP), and low-medium-energy (LME) collimators are also popular in Japan. Single-photon emission computed tomography (SPECT) imaging also allows 360° or 180° rotation and it can score defects similar to perfusion defects (7,8). The Japanese Society Nuclear Medicine working group created normal early / late, 180° / 360° and gender-specific databases (9) that work with any software and are applicable to clinical and research purposes.

Heart-to-mediastinum ratio: ROI setting and stability

The most popular index of cardiac MIBG uptake is early and late HMR. Although the HMR is a simple average count ratio between the heart and mediastinum, the location, size and shape of the ROI results in variability. While the European Association of Nuclear Medicine Cardiovascular Committee and the European Council of Nuclear Cardiology defines no specific ROI size (4), Fig. 1 compares the ROI settings recommended by the American Society of Nuclear Cardiology (10) and the semiautomatic smartMIBG software used in Japan (5). This software needs only to point towards the center of the heart, and then a circular ROI on the heart and a rectangular ROI on the upper mediastinum are automatically determined. The mediastinal ROI was set at 10% of the body width and 30% of the height from the center of the heart to the upper border of the mediastinum. The optimal mediastinal region was automatically searched vertically to determine the minimal count on the mediastinum. The HMR remains relatively stable for three or four hours, when late images can be acquired (11, 12). A washout rate (WR) can also be calculated using the formula:

\[
\text{WR (\%)} = \frac{\text{early heart count-late heart count}}{\text{early heart count}} \times 100.
\]

The mediastinal count in this formula is usually subtracted from the heart count as the background, and a \(^{123}\text{I}\) (half-life, 13 h) decay is corrected using a decay factor at three to four hours after the initial image acquisition (correction factor of \(\times 1.17\) and \(\times 1.24\), respectively). Although ROI settings are generally considered reproducible (13), the semiautomatic algorithm significantly improved inter- and intra-observer variations (5).

Calibration phantom to overcome camera-collimator differences

Differences between collimators, particularly LE and ME types, cause variations in HMR measurements (14). A cross-calibration phantom was therefore designed to calibrate HMR measured in various hospitals (15,16). Two fixed HMRs were calculated from anterior and posterior planar images. Since the mathematically calculated HMR is known, two data points are obtained, and a linear regression line that passes through the coordinate (1,1) for the measured versus the reference HMR is calculated. The slope of this regression line is defined as a conversion coefficient (CC), and it is unique for each scinticamera-collimator system. We proposed unifying the HMR to the ME type of collimator, which conforms to European recommendations and is popular worldwide (4).

Since the average CC of ME general-purpose collimators is 0.88 (16), any institutional HMR (HMRi) can be standardized to MEGP-collimator condition (HMRstd) using the formula:

\[
\text{HMRstd} = \frac{0.88}{\text{CCi}} \times (\text{HMRi}-1)+1,
\]

where CC is the conversion coefficient of an institutional camera-collimator system. We proposed unifying the HMR to the ME type of collimator, which conforms to European recommendations and is popular worldwide (4).

The average CC of typical collimators are: 0.55, 0.65, 0.62 or 0.75, 0.83, 0.88 and 0.95 for LEHR, LEGP, extended LE general-purpose (ELEGP) with two types depending on camera, LME, MEGP and ME low penetration (MELP) collimators, respectively (16). However, the CC of the same LEHR collimator that was commonly used in the multicenter ADMIRE-HF study might have significantly varied depending on septal thickness, the size and length of the hole, as well as the camera crystals (17).

The HMR can also be calculated from images acquired using single-photon emission computed tomography (SPECT), for which custom-designed software is required to sum myocardial photon counts and set appropriate background regions (18). The SPECT system with solid-state detector technology, D-SPECT, has high sensitivity and resolution but it is unsuitable for obtaining planar anterior images. However, planograms comparable with planar anterior images can be reconstructed (19). Based on calculations similar to those used in conventional HMR, D-SPECT and Anger SPECT findings closely correlate. Moreover, converting the Anger HMR to the standardized HMR with a CC of 0.88, renders the HMR essentially identical, indicating that HMR derived from conventional planar images and images acquired using new SPECT cameras can be integrated (20).

Difference in collimator types did not affect WR values significantly. When the JSNM working group MIBG database...
LEGP and ME collimators. When the estimated average was
of 0.6) was 1.68 (23), which was again converted to 2.0 with
(using both LEHR and LEGP collimators with an average CC
analyses of a pooled database from six Japanese hospitals
the standard ME-collimator. The threshold determined from
institutions (17). When the value is converted using an average
death and lethal arrhythmia was 1.6 in the ADMIRE-HF study
in which LEHR collimators were used at all participating
centers (1,17,21) and the HMR in Japanese prognostic studies
significantly differ among several MIBG studies at various
centers (1,17,21) and the HMR in Japanese prognostic studies
is slightly higher. Although the background differs among
studies according to the baseline status of patients, differences
in collimators might have been involved. Japanese vendors
have since attempted to optimize collimator design for the
higher-energy photons emitted by 123I radiopharmaceuticals
that are popular in cardiac and brain studies.

For example, the optimal threshold for predicting cardiac
death and lethal arrhythmia was 1.6 in the ADMIRE-HF study
in which LEHR collimators were used at all participating
institutions (17). When the value is converted using an average
CC of 0.55, the threshold HMR can be interpreted as 2.0 with
the standard ME collimator. Nakata et al. determined that a
threshold of 1.74 for prognosis of CHF using an LEGP
collimator (CC=0.65) (22), and it can be converted to 2.0 with
the standard ME-collimator. The threshold determined from
analyses of a pooled database from six Japanese hospitals
(using both LEHR and LEGP collimators with an average CC
of 0.6) was 1.68 (23), which was again converted to 2.0 with
the standard ME collimator. Agostini et al. summarized
European databases using a threshold of 1.75 (24) and LEHR,
LEGP and ME collimators. When the estimated average was
0.6 based on the weighted average of their data, the corrected
value was around 2.0.

To better understand the value of 123I-MIBG imaging, all
threshold values in the literature need to be re-evaluated,
regardless of the baseline status of patients and study purpose.
Larger databases for prognostic studies could be generated
after original databases are created.

Application of standardized MIBG HMR to mortality risk model
Accumulating clinical evidence shows that the MIBG HMR
is useful for predicting lethal cardiac events. However, actual
risk for cardiac death cannot be evaluated by HMR alone.
Multicenter studies such as ADMIRE-HF, a European MIBG
meta-analysis, and a Japanese pooled database analysis
showed that around 1.6-1.7 is the late HMR threshold for
predicting a poor prognosis as alluded to above. Therefore, a
cardiologist receiving for example, an MIBG report of a
patient with CHF including HMR=1.55, would predict a poor
prognosis. However, the significance of what the MIBG
findings indicate is not exactly intuitive because cardiologists
judge individual patient risk after considering several factors
such as symptoms of heart failure, age, basic cardiac function,
arrhythmia, medications, complications with diabetes mellitus
and chronic kidney disease, and cardiac devices such as
implantable cardioverter defibrillators. We therefore created
cardiac mortality risk models based on multicenter pooled
databases and used multivariate analysis to select the five most
potent variables, namely, New York Heart Association
(NYHA) functional class, age, sex, left ventricular ejection
fraction (LVEF) and 123I-MIBG HMR (6). To establish
relatively short and long-term (two and five year, respectively)
risk models, we subsequently selected the categorical
variables of NYHA class (I-II and III-IV), age (<65 and ≥65
years), LVEF (<35%, 35%-50%, >50%) and HMR (<1.40,
1.40-1.69, 1.70-1.99 and ≥2.00) as variables (25). This model
was applicable to both ischemic and non-ischemic etiologies
of CHF. It can also be used for evaluating effectiveness of
pharmacological treatment and personalized patient care in
patients with heart failure (26,27).

That Japanese cohort study proceeded using LEHR and
LEGP collimators between 1990 and 2009, and the cardiac
risk model was generated using the averaged values of LE
collimators. However, current 123I-MIBG studies in Japan use
LEGp, ELEGP, and LME collimators, which result in a higher
HMR compared with that generated during the 1990s.
Although our recommendation is standardization to the ME
collimator with a conversion coefficient of 0.88, all HMRs
should be converted to the LE collimator (average CC of 0.6 in
the Japanese pooled database) in the internal calculation to
apply this risk model to a current study. When risk models

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**Fig. 2** Conversion coefficients for various collimators
Average and standard deviation are shown for each collimator
(n = 62) was analyzed (9), WRs for LE and ME/LME
collimators were 13 ± 7% and 14 ± 10%, respectively (p=n.s.),
and 13 ± 8% as a whole. To utilize WR, however, application
of the time-decay correction (usually 3 to 5 hours),
background correction and time decay correction between
early and late imaging should be unified among studies.

**Application of standardization in the literature**
Normal values and thresholds for predicting cardiac events
significantly differ among several MIBG studies at various
centers (1,17,21) and the HMR in Japanese prognostic studies
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the Japanese pooled database) in the internal calculation to
apply this risk model to a current study. When risk models
were preliminarily compared between calculations based on CCs of 0.6 and 0.88, the final predicted mortality risk was nearly identical. Therefore, by combining conversion formulae among collimators, the five-year cardiac mortality risk chart became applicable to any type of collimator as shown in Fig. 3.

Conclusion
Standardization of \(^{123}\text{I}\)-MIBG parameters, in particular HMR, plays a pivotal role in the diagnosis, treatment and prognostic estimation of CHF, whereas quantitation methods based on SPECT might progress. Cardiac mortality risk models could be more flexibly applied to various stages of CHF at any institution using standardized MIBG parameters.

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Conflicts of interest
K. Nakajima participates in a collaborative research project with FUJIFILM RI Pharma Co. Ltd. to develop smartMIBG software.

Reprint requests and correspondence:
Kenichi Nakajima, MD, PhD
Department of Nuclear Medicine, Kanazawa University Hospital, 13-1 Takara-machi, Kanazawa, Japan 920-8641
E-mail: nakajima@med.kanazawa-u.ac.jp


