OVERVIEW

This report summarizes the development, administration, and scoring of the nuclear cardiology specialty examination offered by the Nuclear Medicine Technology Certification Board (NMTCB) at the Annual Meeting of the Society of Nuclear Medicine in Toronto, Canada, on June 23, 2001.

During the NMTCB meeting in the fall of 2000, the Board approved the development and administration of an advanced practice nuclear cardiology examination in response to a formal request from the Society of Nuclear Medicine—Technologist Section and the Nuclear Cardiology Committee. Many years of experience in developing and setting standards for the entry-level nuclear medicine certification examination aided the NMTCB in taking on this new responsibility. From its existing members, the NMTCB formed an ad hoc nuclear cardiology specialty examination committee composed of technologists, educators, a pharmaceutical specialist, a physician, and psychometric staff. The task of writing the examination began immediately.

PURPOSE AND REQUIREMENTS

The primary purpose of this specialty examination was to assess the knowledge beyond the entry level of an experienced nuclear cardiology technologist. Eligible candidates who took the examination were certified in entry-level nuclear medicine technology and had a minimum of 2 y full-time clinical experience in nuclear cardiology. Holding an active NMTCB, RT(N), or CAMRT certification fulfilled the entry-level certification requirement. The NMTCB office in Atlanta processed all applications, including verification of candidates’ entry-level certificate and 2-y work experience. The certification received by candidates after successfully passing the examination is valid up to 7 y from the examination date and will expire thereafter unless the individual chooses to re-certify. Re-taking and successfully passing the specialty examination could accomplish re-certification. Upon successful completion of the examination, the technologists were granted the right to use the title Nuclear Cardiology Technologist (NCT).

EXAMINATION DEVELOPMENT

A subject-matter expert (SME) group was involved in identifying critical abilities related to the practice and in defining test specifications. This SME group included 5 members of the ad hoc committee and 5 professionals practicing in the nuclear cardiology field who came from different regions of the country and from different work settings for a total of 10 members. The Guidelines for Technologist Training in Nuclear Cardiology (1), developed by the technologist committee of the American Society of Nuclear Cardiology, a leading professional organization in the field, was used to verify and refine the content domain for the examination. The SME group evaluated relevant documents and identified the examination content outlines and assigned a percentage to each piece of content by consensus. The SME group also determined the test length to be 100 multiple-choice items with 4 options and identified an appropriate testing time to be 2 h. The ad hoc committee members wrote most of the test items. The SMEs and other members of the Board completed the pilot test. After the pilot test, the group worked together to finalize the content outlines for the test, percentage of the total number of items to be used from each content area, as well as test items to be used on the test. For security reasons 2 test forms, A and B, were developed. By scrambling the item order of form A, form B was created. The content outline with the percentage specified in parentheses from each content area is presented below. For a detailed list of the content outline, procedures, and pharmaceuticals list see the Appendix.

Content Outline

I. Instrumentation, procedures, and processing: myocardial perfusion imaging (38%); and equilibrium radionuclide angiocardiography, first pass, and shunts (12%)

II. Anatomy, physiology, and pathology (10%)
III. Radiopharmaceuticals and interventional drugs (15%)

IV. Nonpharmacologic (exercise) stress testing (15%)

V. Patient care (10%)

EXAMINATION ADMINISTRATION

A total of 59 candidates were registered to take the first nuclear cardiology examination in Toronto. Only 58 candidates completed the test, 36 in the morning (Form A) and 22 in the afternoon (Form B). Candidates were checked into the testing room by verifying their identification with the roster prepared by the Board office. The candidates were not allowed to choose their own seats but were assigned seats by the proctor. The testing was administered at the Fairmont Royal York Hotel, Toronto, Canada. The examination room provided adequate writing surface, uncrowded seating, good lighting, comfortable temperature, and freedom from distraction. The test included 100 scored and 21 pretest items. All graphics that were part of the multiple-choice items of the test were collated in a separate booklet. Candidates were given a total time of 2 h to complete the test. The proctor began the session by reading a standard set of examination instructions. Candidates were required to sign a form for maintaining the confidentiality of the test questions. They were given notification of the remaining testing time at 30, 10, and 5 min from the end of the 2-h allotment. The morning group of candidates was not dismissed after completing the test, but was escorted to a different room until all of the candidates in the afternoon session were checked in. All candidates were encouraged to submit written comments about the test on the back of the examination booklet. In each session only 5 candidates were still in the examination room when the proctor made the final, 5-min-remaining announcement of the examination time.

TEST SCORING

Before scoring, all answer sheets were verified as to their accuracy. Answer sheets from the 2 sessions were machine-scored separately. Multiple response and blank responses from candidates’ answer sheets recorded by the machine were verified. Upon verification it was found that the machine recorded multiple responses for incompletely erased bubble marks and blank responses for lightly bubbled responses. These errors were corrected before generating a preliminary item analysis report. The analysis report was conducted to detect key errors and items that were not performing properly. Members of the ad hoc committee reviewed the item analysis report and comments received from candidates. As a result of the review, 6 items were removed from scoring and 3 items were scored with a double key. All candidates were graded on 94 test items. The passing cutoff score for the test was set to be 75 on the scaled score. The reliability (KR-20) of test forms A and B were 0.86 and 0.83, respectively.

Table 1 gives the number of items based on the range of difficulty and discrimination indexes on both test form A and B. The histogram of the examinees’ scaled scores is presented in Figure 1.

TABLE 1  Number of Items Grouped by Discrimination and Difficulty Indexes

<table>
<thead>
<tr>
<th>Range</th>
<th>Discrimination index count</th>
<th>Difficulty index count</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Form A</td>
<td>Form B</td>
</tr>
<tr>
<td>1.00</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>0.99–0.90</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>0.89–0.80</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>0.79–0.70</td>
<td>---</td>
<td>1</td>
</tr>
<tr>
<td>0.69–0.60</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>0.59–0.50</td>
<td>2</td>
<td>6</td>
</tr>
<tr>
<td>0.49–0.40</td>
<td>10</td>
<td>6</td>
</tr>
<tr>
<td>0.39–0.30</td>
<td>16</td>
<td>10</td>
</tr>
<tr>
<td>0.29–0.20</td>
<td>24</td>
<td>22</td>
</tr>
<tr>
<td>0.19–0.10</td>
<td>21</td>
<td>17</td>
</tr>
<tr>
<td>0.09–0.00</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>&lt;0.00</td>
<td>7</td>
<td>11</td>
</tr>
</tbody>
</table>

In summary, the examination appears to exhibit appropriate statistics for a credentialing examination. The job
analysis model and cutoff-score study described in this report help to substantiate the validity of the resulting examination scores.

APPENDIX

Content Outline

I. Instrumentation, procedures, and processing: myocardial perfusion imaging (~38%); and equilibrium radionuclide angiocardiography, first pass, shunts (~12%)

A. Acquisition
   1. Patient preparation, indications, contraindications, sequencing of procedures
   2. Injection techniques
   3. Acquisition protocols
      a. Patient positioning
      b. Stopping parameters
      c. Matrix size
      d. SPECT parameters
      e. Gating parameters
         i. Frame rate
         ii. Acceptance window
      f. Collimator

B. Processing
   1. Cineangiograms
   2. Ejection fraction determination
   3. Functional images
   4. Heart–lung ratio
   5. Image manipulation techniques
   6. Image filtering
   7. Polar-plot analysis
   8. Wall–motion analysis
   9. Time–activity curves

C. Quality control
   1. Image assessment techniques
   2. Quality assessment of filtering techniques

D. Artifacts
   1. Motion
   2. Center-of-rotation correction errors
   3. Attenuation
   4. Radiopharmaceutical distribution
   5. Artifacts created by processing techniques

II. Anatomy, physiology, and pathology (~10%)

A. Heart chambers
B. Cardiac electrophysiology
   1. Conduction pathways
   2. Normal electrocardiogram
C. Coronary artery distribution
D. Heart valves and great vessels
E. Normal and abnormal physiologic responses to stress
F. Cardiac pathologies
   1. Coronary artery disease
   2. Cardiomyopathies
   3. Myocardial ischemia, infarction, hibernation, and stunning
   4. Valvular diseases and effects on the heart
   5. Congenital cardiac anomalies

III. Radiopharmaceuticals and interventional drugs (~15%)

A. Radiopharmaceuticals
   1. Indications
   2. Dosages
   3. Radiopharmaceutical problems

B. Interventional drugs
   1. Types and dosages
   2. Indications

FIGURE 1. Histogram of NMTCB nuclear cardiology examinees’ scaled scores.
3. Pharmacologic stress protocols
4. Contraindications and medication interactions

IV. Non-pharmacologic (exercise) stress testing (~15%)
A. Contraindications to exercise stress testing
B. Physiologic measures of exercise capacity and performance
C. Electrocardiogram (ECG) acquisition
D. Treadmill tower operation
E. Patient monitoring
F. Bicycle and isometric exercise protocols
G. Patient assessment and monitoring
H. Endpoints

V. Patient Care (~10%)
A. Answering patient questions
   1. Risks of nuclear medicine procedures
   2. Comparison with correlative imaging techniques
B. ECGs
   1. Patient preparation, electrode placement, and leads
   2. Rate calculation
   3. Normal and abnormal rhythms
   4. Heart blocks
   5. Indicators of ischemia and infarction
C. Emergency care
   1. CPR
   2. Emergency medications
   3. Diabetic complications

Procedures List
I. Myocardial perfusion study
   A. Treadmill, bicycle, or isometric exercise
   B. Pharmacologic stress
   C. Planar imaging
   D. SPECT
   E. Gated SPECT
   F. PET
II. Equilibrium radionuclide angiocardiogram
   A. Rest
   B. Exercise
   C. SPECT
III. Left-to-right cardiac shunt study
IV. Right-to-left cardiac shunt study
V. First-pass study using a multicroystal system
VI. Gated first-pass study

Pharmaceuticals List
1. $^{123}$I-MIBG
2. $^{99m}$Tc-sestamibi
3. $^{99m}$Tc-tetrofosmin
4. $^{201}$Tl-thallous chloride
5. $^{99m}$Tc-labeled red blood cells
   a. In Vivo labeling
   b. Modified In Vivo and In Vitro labeling
   c. Ultra Tag® (Mallinckrodt, St. Louis, MO)
6. $^{18}$F-FDG
7. $^{13}$N ammonia
8. $^{15}$O water
9. $^{82}$Rb chloride
10. $^{11}$C fatty acid
11. Adenosine
12. Dipyridamole
13. Aminophylline
14. Dobutamine
15. Esmolol hydrochloride
16. Acetyl salicylic acid
17. Anticoagulants
18. Antiarrhythmics
19. Calcium channel blockers
20. Angiotensin converting enzyme (ACE) inhibitors
21. Nitrates
22. Beta blockers
23. Cholesterol-lowering drugs
24. Diuretics
25. Angiotensin receptor blockers
26. Digoxin

REFERENCE