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Abstract

Coronary artery disease is a leading cause of morbidity and mortality in developed countries. According to a Center for Disease Control report, one out of four deaths is attributed to coronary artery disease. It costs the United States human lives, productivity, and more than 100 billion dollars each year. Due to increased incidence in both men and women and all ethnicities, risk stratification of patients at risk for developing myocardial infarction and death is of paramount importance. Various tests are available for diagnosis and prognosis in coronary heart disease such as exercise treadmill testing, coronary calcium scoring, dobutamine stress echocardiography, exercise, dipyridamole, adenosine or dobutamine stress nuclear myocardial perfusion imaging (MPI), and dobutamine or adenosine stress cardiac magnetic resonance imaging. Since 2008 a new vasodilator, regadenoson (REG), has become available and is now widely used for nuclear perfusion imaging. Pharmacologic stress testing challenges the coronary flow reserve to evaluate the hyperemic capacity of the heart, which can be impaired in significant epicardial stenosis or microvascular dysfunction. In the presence of either of these conditions, ischemia induced by hyperemia manifests as wall motion abnormalities on echocardiography or as perfusion defects in nuclear perfusion imaging.

REG is a selective adenosine $A_2A$ receptor agonist, and due to its targeted coronary vasodilator properties and bolus administration of a standard dose in all patients, it has rapidly gained popularity as the preferred MPI stress agent. In this chapter we will review the basis of pharmacologic vasodilator stress imaging starting with a brief discussion of the various adenosine receptors and their function, the structure and mechanism of action of REG, and its development and approval. It will be compared with other myocardial perfusion pharmacologic stress agents like adenosine and
dipyridamole in terms of safety, efficacy, and side effect profile. We will also address the utility of REG in special situations like renal disease, chronic obstructive pulmonary disease, heart transplant, left bundle branch block, and paced rhythms. The prognostic value of REG MPI in the general population, its effectiveness with and without exercise, and the emerging applications of REG in other modalities of imaging such as positron emission tomography and stress echocardiography will be discussed.

**Keywords:** Regadenoson, single photon emission computed tomography, positron emission tomography, stress echocardiography, fractional flow reserve, coronary artery disease

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1. Introduction

Cardiovascular disease remains a leading cause of death in the United States. According to a 2009 report by the Center for Disease Control, one out of four deaths is attributable to coronary artery disease (CAD).[1] The increased morbidity and mortality due to CAD poses a huge economic burden. In 2010, CAD alone accounted for over 100 billion dollars in combined direct and indirect (i.e., loss of productivity) costs. This is projected to more than double by 2030.[2] Hence, diagnosing and risk stratifying CAD in its early stages is vital.

Many invasive and noninvasive tests are available to identify patients at high risk of developing CAD. Functional tests include exercise stress testing, exercise or dobutamine stress echocardiography, nuclear myocardial perfusion imaging (MPI) using stress agents such as dipyridamole, adenosine, dobutamine or regadenoson (REG), and vasodilator stress magnetic resonance imaging. Coronary computed tomography angiography (CCTA) and the traditional gold standard, coronary angiography, serve as the two well-established anatomic modalities used for CAD detection. This chapter will focus on REG, the newest of the pharmacologic stress agents, and its applications in myocardial perfusion imaging. It will conclude with a brief overview of some novel applications of REG in cardiology.

2. REG: development, pharmacology, and hemodynamic effects

2.1. Adenosine receptors

Adenosine receptors are located in the myocardium as well as in smooth muscle cells of the coronary arterioles and the bronchial tree. Various subtypes of adenosine receptors exist including $A_1$ receptors found in the atrioventricular node, $A_{2A}$ receptors present in coronary arteriolar smooth muscle, and $A_{2B}$ and $A_3$ receptors located in bronchial smooth muscle. The different locations and functions of these receptors have been pivotal in the development of newer pharmacologic stress agents (Figure 1). Adenosine directly and dipyridamole indirectly act on adenosine 2A ($A_{2A}$) G-protein-coupled receptors found on the cell membrane of coronary
arteriolar smooth muscle cells. However, both are nonselective and also activate the other adenosine receptor subtypes causing frequent clinically important side effects (e.g., atrioventricular block due to $A_1$ activation and bronchoconstriction due to $A_{2B}$ and $A_3$ receptor activation) as well as other less serious but often unpleasant side effects. In contrast, REG exerts its effect selectively on $A_{2A}$ receptors achieving the coronary dilatation necessary to perform MPI studies while keeping side effects to a minimum.

![Diagram of adenosine receptors and pharmacologic agents](http://dx.doi.org/10.5772/61154)  

Note: Springer and J Nucl Cardiol, 17, 2010, 494-497, The emerging role of the selective A2A agonist in pharmacologic stress testing, Gemignani AS, Abbot BG, Figure 1. With kind permission from Springer Science and Business Media

**Figure 1.** Types of adenosine receptors, their functions, and activation/inhibition by various pharmacologic agents.

### 2.2. Development and approval of REG

Cardiac stress testing is able to identify as well as risk stratify individuals who are at risk for CAD. Vasodilator stress testing challenges the coronary flow reserve in order to evaluate the hyperemic capacity of the heart, which can be impaired in significant epicardial stenosis or microvascular disease and lead to transient ischemia. Ischemic changes manifest either as perfusion or wall motion abnormalities depending on the imaging modality used. The currently available pharmacologic stress agents with primarily vasodilator function are dipyridamole, adenosine, and REG. While dobutamine also vasodilates, it mainly stresses the heart via its positive inotropic and chronotropic effects.

An ideal cardiac stress agent should cause short-lived but maximal coronary vasodilatation. Both of these can be achieved if the stress agent has low affinity for its receptor and the target tissue has many adenosine receptors. The coronary arterial tree has an abundance of $A_{2A}$
receptors of which only a fraction needs to be activated to elicit the desired coronary vasodilation and produce maximal coronary hyperemia. Given the nonspecific nature of adenosine receptor stimulation by adenosine and dipyridamole leading to undesired side effects, the need existed for the development of an $A_{2A}$-selective agent largely devoid of significant side effects such as bronchospasm and atroventricular conduction block. REG (code name CVT 3146) was identified as an agent with $A_{2A}$ selectivity yet with a low affinity for $A_{2A}$ receptors, meaning it dissociates quickly after eliciting maximal coronary vasodilation, thus causing adequate coronary hyperemia for a short period of time. REG underwent preclinical and subsequently randomized clinical studies showing non-inferiority compared to the commonly used vasodilator adenosine. This led to its approval by the Food and Drug Administration in 2008. It is marketed by Astellas Pharma US Inc. under the trade name Lexiscan® in the United States as a cardiac stress agent for MPI studies in patients who are unable to exercise. Following REG administration, coronary hyperemia occurs for approximately 2–5 min, which is adequate for radionuclide uptake and makes it possible to perform stress testing using a single bolus injection.[3]

2.3. Pharmacology and pharmacokinetics of REG

REG is a 2-[[N-1-(4-N-methylcarboxamidopyrazolyl)] adenosine derivative. It is prepared by condensing ethoxycarbonylmalondialdehyde with 2-hydrazinoadenosine in a 1:1 mixture of ethanoic acid and methanol. The resulting ester is then converted directly by aminolysis with methylamine to the amide REG (Figure 2). Alternatively, REG can be prepared from 2-chloro or 2-iodo adenosine derivatives. The amide links at the 4-position of the $N$-pyrazolyl, which has both lipophilic and hydrophilic substituents lending the drug greater affinity for the adenosine $2A$ receptor than the other adenosine receptor subtypes.

![Figure 2. The molecular structure of REG (CVT-3146; (1-[9-(4S, 2R, 3R, 5R)-3,4-dihydroxy-5-(hydroxymethyl)oxolan-2-yl]-6aminopurin-2-yl]pyrazol-4-yl]-N-methylcarboxamide).](image)

It is usually given as a single 400-μg (5 mL) intravenous bolus after which it immediately distributes throughout the body. No weight-based dose adjustment is necessary. REG then undergoes three phases of elimination. The first is the phase of maximal coronary hyperemia.
lasting 2–4 min.[4] The second phase lasts 15–30 min with profound effect on heart rate and blood pressure, and the third phase, which lasts for 33–108 min, is clinically nonsignificant.[5] Much about REG’s metabolism remains unknown; however, its excretion is both renal and hepatic. The kidneys remove approximately 60% via tubular secretion, while the liver excretes around 40% of the drug unmetabolized into the bile.

2.4. Hemodynamic effects of REG

As a coronary vasodilator REG is shown to cause tachycardia and changes in blood pressure (both increase and decrease). Trochu et al.[6] showed in animal studies that while adenosine increased left ventricular (LV) systolic pressure, REG did not to any significant degree, and that LV contractility measured by dP/dT increased by 39±7% with REG and 29±7% with adenosine. The ADVANCE MPI studies[7] have shown that the decrease in systolic and diastolic blood pressures (BP) was similar between REG and adenosine (systolic BP drop 14±13 mmHg vs. 13±14 mmHg, P = ns; diastolic BP drop 10±8 mmHg vs. 10±8 mmHg, P = ns). Both drugs increase the heart rate; however, REG more significantly than adenosine (25±11bpm vs. 20±10bpm, P < 0.001).

The increase in heart rate with REG is mainly due to direct sympathetic excitation and less so from a baroreceptor reflex induced tachycardia. Dhalla et al.[8] has also suggested that an A_2A receptor mediated sinus tachycardia can occur with REG. A blunted heart rate acceleration with both REG and adenosine has also been observed in studies with diabetic patients and is felt to be related to sympathetic denervation.[9]

2.5. Side effect profile of REG

Like other vasodilators, REG is associated with many minor and a few major (albeit to a lesser extent than older vasodilators) side effects of which clinicians need to be aware.[10] Transient side effects included nausea (6%), abdominal pain (5%), headache (26%), and chest tightness (13%). In the randomized studies evaluating REG prior to its FDA approval, atrioventricular (AV) block incidence was <1% with no instances of advanced AV block or asystole in the ADVANCE MPI 3 studies. However, post marketing surveillance has highlighted rare major adverse reactions related to REG such as acute myocardial infarction,[11, 12] atrioventricular block, and asystole.[13] Thus, REG, despite its A_2A selectivity, should not be used in patients with greater than the first-degree AV block unless they have a backup pacemaker. Furthermore, cases of syncope[14] and seizures[15] have also been reported following REG administration. Although aminophylline is used for reversal of many REG-induced side effects, it should not be used in the setting of seizures following REG injection as it lowers the seizure threshold. Instead, standard antiseizure therapy with benzodiazepines and agents such as phenytoin should be used.

2.6. Effect of caffeine on REG and clinical implications

Caffeine is an A_2A receptor antagonist (Figure 1). Hence, it has the potential to attenuate the hyperemic response, which occurs after vasodilator administration. This is a well-known
problem with adenosine and dipyridamole, both of which require abstinence from caffeinated products for at least 24 h prior to stress testing. However, the REG package insert specifies withholding caffeinated products for only 12 h prior to testing. Preclinical animal studies suggested that caffeine attenuated the duration of REG-induced coronary hyperemia in dogs. Subsequent human studies evaluating myocardial blood flow in 41 healthy volunteers using REG with PET imaging showed that moderate caffeine consumption may not interfere with REG-induced coronary hyperemia. Thus, conflicting evidence existed regarding the effect of caffeine on REG stress testing until a multicenter randomized trial on this subject was performed in 2014.

Tejani et al.[18] studied the effects of caffeine on the diagnostic accuracy of REG single proton emissions computed tomography (SPECT) MPI in 207 subjects with documented coronary artery disease on an initial rest-REG SPECT MPI sequence. A third set of SPECT images was acquired in all patients following randomization to two different caffeine doses (200 and 400 mg) or placebo. Previously noted reversible defects were attenuated in patients who consumed both doses of caffeine at least 90 min prior to REG administration, thus diminishing the diagnostic accuracy of the study. There was no difference in adverse effects between the three groups.[18] Current American Society of Nuclear Cardiology (ASNC) guidelines recommend that patients refrain from caffeine consumption for at least 12 h before REG stress testing.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Regadenoson</th>
<th>Adenosine</th>
<th>Dipyridamole</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brand name</td>
<td>Lexiscan®</td>
<td>Adenocard®/Adenoscan®</td>
<td>Persantine®</td>
</tr>
<tr>
<td>Indication</td>
<td>Pharmacologic stress agent in MPI</td>
<td>Treatment of paroxysmal supraventricular tachycardia, pharmacologic stress agent in MPI</td>
<td>Oral—antithrombotic along with warfarin/aspirin. Intravenous—pharmacologic stress agent in MPI</td>
</tr>
<tr>
<td>Mechanism of action</td>
<td>Increases coronary flow reserve (CFR) via selective A₁₃ adenosine receptor agonism</td>
<td>Nonselective adenosine agonist on A₁, A₂A, A₂B, and A₃ receptors. Increases coronary flow reserve (CFR) via A₁ receptor activation.</td>
<td>Increases availability of adenosine by inhibiting adenosine deaminase, which prevents adenosine’s breakdown</td>
</tr>
<tr>
<td>Potency</td>
<td>10 times more potent than adenosine</td>
<td>Less potent</td>
<td>Less potent</td>
</tr>
<tr>
<td>Distribution in body</td>
<td>11.5 L</td>
<td>Unknown</td>
<td>2–3 L</td>
</tr>
<tr>
<td>Metabolism</td>
<td>Unknown</td>
<td>In blood and tissue, metabolized by adenosine deaminase into inosine and then adenosine monophosphate and hypoxanthine</td>
<td>Hepatic</td>
</tr>
<tr>
<td>Time to peak</td>
<td>1–4 min</td>
<td>30 s</td>
<td>2–2.5 h</td>
</tr>
</tbody>
</table>
Variable | Regadenoson | Adenosine | Dipyridamole
--- | --- | --- | ---
Half-life | Triphasic | First phase = 2–4 min | Second phase = 15–30 min | Third phase = 33–108 min | <10 s | 30–45 min
Administration | Bolus | Infusion | Infusion
Dose | 400 μg | 140 μg/kg/min | 0.14 mg/kg/min
Duration of infusion | 10-20 s bolus | 6 min continuous infusion | 4 min continuous infusion
Excretion | 57% of drug excreted unchanged in urine via tubular secretion | Cellular uptake | Conjugated by glucuronide and unchanged drug excreted in feces
Safety in pregnancy | Risk cannot be ruled out (Category C) | Risk cannot be ruled out (Category C) | No evidence of human risk in controlled studies (Category B)
Common side effects | Headache 26%, flushing 16%, dyspnea 28%, hypotension 2% | Headache 21%, flushing 35%, dyspnea 19%, hypotension 3% | Headache 12%, flushing 3.4%, dyspnea 2.6%, hypotension 5%
IV tubing | Not needed: only Hep-lock | Needed | Needed
Protocol completion time with radiotracer | Less than 1 min | 4–6 min | 6–8 min

Table 1. Comparison of the three commonly used vasodilator agents

Compared with the other two agents, REG is more potent, causes more selective coronary vasodilatation, can be injected in a single bolus without weight-based adjustments, and produces SPECT images comparable to adenosine and dipyridamole.

3. REG SPECT MPI in detection of coronary artery disease

3.1. Comparison to adenosine

In a multicenter phase 2 study, REG was tested in 36 patients undergoing SPECT MPI at bolus doses of 400 and 500 μg. Patients with heart transplantation, left bundle branch block, ventricular pacemaker, and low ejection fraction (14 patients) were excluded. This study showed a higher rate of detecting reversible perfusion defects with the lower dose of REG (89% for 400 μg) than with the higher dose (76% for 500 μg).[19] Subsequently, two phase 3 double-blinded, randomized, multicenter trials (ADVANCE-MPI 1 and ADVANCE-MPI 2) demonstrated non-inferiority of REG SPECT MPI to adenosine SPECT MPI. The ADVANCE-MPI 2 trial included 54 sites and 784 patients undergoing clinically indicated adenosine MPI who were blindly randomized 4 weeks later to a second MPI study with REG (n = 495) or adenosine (n = 260) in a 2:1 ratio. Study images were reported in a blinded fashion by three nuclear
cardiology experts unaware of any patient data. The primary aim of the study was to show
the strength of agreement between sequential adenosine and REG images, and the non-
inferiority of the adenosine-REG sequence to the adenosine-adenosine sequence for consistently detecting reversible perfusion defects. The investigators demonstrated that the overall agreement was not statistically different between sequential adenosine–adenosine images (0.64 ± 0.04) compared to adenosine–REG images (0.63 ± 0.03). Furthermore, there was no significant difference in image quality between the two stress agents, and the patient tolerability questionnaire favored REG in this study. In a subsequent quantitative analysis of the ADVANCE-MPI 2 study, investigators showed that the total perfusion defect size, ischemic perfusion defect size, ejection fraction, and LV volume estimation was similar between REG and adenosine.[20] Thus, cumulative evidence collected from over 2000 patients in these pivotal phase 3 trials demonstrated the non-inferiority of REG to adenosine in SPECT MPI,[7] as well as the effects of age, gender, obesity, and diabetes on the efficacy and safety of REG[21] leading to its approval for clinical use.

3.2. REG in special populations

3.2.1. Renal disease

The predominantly renal excretion of REG (60% of the drug) raises concern for its safety in chronic kidney disease and end stage renal disease patients, including those on dialysis. To date, two major studies and one prognostic study have shown that REG is not associated with any major adverse events in this group.

Ananthasubramaniam et al.[22] conducted a randomized, double-blinded, placebo-controlled multicenter trial to evaluate the safety and tolerability of REG in 432 patients with stage 3 (glomerular filtration rate (GFR) 30–59 mL/min/1.73 m²) and 72 patients with stage 4 (GFR 15–29 mL/min/1.73 m²) chronic kidney disease. There were no major adverse events within 24 h of REG injection in the intervention group. Minor adverse effects like headache, dyspnea, chest discomfort, nausea, flushing, and dizziness were more common in the REG group than in the placebo group.

Doukky et al.[23] studied 146 ESRD patients undergoing REG stress testing, which included 131 patients on hemodialysis, 12 patients on peritoneal dialysis, and two not on any dialysis. These were compared with 97 control patients with GFR ≥30 mL/min. The primary end point of the study was patient reported side effects within 24 h following REG administration. There were no statistically significant differences in adverse effects between the groups. Interestingly, end stage renal disease patients tolerated REG stress better than the control group and expressed their willingness to take the test again (117/131 (80%) vs. 63/97 (65%), P = 0.001).[23]

3.2.2. Asthma and COPD

Adenosine A₂A and 3 receptors are located in bronchial smooth muscle cells which, when activated, can lead to bronchoconstriction (Figure 1). Although REG is a selective A₂A receptor
agonist, there is a concern related to its use in patients with asthma and chronic obstructive pulmonary disease (COPD).

More than six studies have been performed to evaluate the safety of REG in this population specifically looking at respiratory symptoms, spirometry parameters, hemodynamic response, and major adverse events. The combined population of these five prospective studies and one retrospective study comprised 686 COPD patients and 695 asthmatics.[24] Respiratory parameters like FEV1, FVC, FEV1/FVC ratio, and patient-reported symptoms were monitored in most of these studies. All showed that REG is safe in COPD and asthmatics. Dyspnea was reported more frequently in COPD and asthmatics, but no significant decline in spirometry measurements occurred among these patients in two double-blinded studies.[25, 26] Of particular note, Kwon et al.[27] demonstrated that patients who underwent low-level exercise in conjunction with REG stress reported fewer respiratory symptoms than those who did not exercise following REG administration.

3.2.3. Pacemaker and left bundle branch block

In patients with left bundle branch block (LBBB), pacemaker, or intrinsic conduction disease, the increased heart rate caused by either exercise, or dobutamine can lead to false-positive septal perfusion defects. This is due to a tachycardia-induced decrease in diastolic perfusion in an already asynchronously activated septum. Multiple studies have compared adenosine and exercise stress tests in these patients. Caner et al.[28–30] showed that dobutamine stress testing is associated with higher false positives in LBBB patients, and similar results were observed in pacemaker patients as well.

The ability of REG to identify perfusion defects in this population was studied by Thomas et al.[31] In their sub-analysis of the ADVANCE MPI 1 and 2 trials, where all 2015 subjects underwent SPECT MPI with adenosine followed by SPECT MPI with either REG or adenosine, 64 patients with LBBB and 93 with pacemakers were identified. Hemodynamic changes, visually assessed summed difference scores (SDS), and quantitative perfusion defects in the LAD territory and septum were compared between REG MPI and adenosine MPI. The study showed that although REG led to a significant increase in heart rate compared with adenosine, it did not cause or exaggerate perfusion defects in the LAD or septal territories either by SDS or quantitative assessment.[31]

3.2.4. Orthotopic heart transplant patients

Orthotopic heart transplant (OHT) patients have a higher incidence of AV block due to denervation supersensitivity. Hence, OHT patients who undergo MPI studies are at increased risk for developing high-grade AV block. Few studies have evaluated the role of MPI in diagnosing cardiac allograft vasculopathy in these patients.

In a retrospective analysis, Al-Mallah et al.[32] identified 102 OHT patients who underwent adenosine MPI and compared them with 204 control patients for heart rate, blood pressure changes, and occurrence of AV block. A threefold increase in the incidence of high-grade AV block (Mobitz type II and third degree) was seen in OHT patients vs. controls. Symptomatic
bradyarrhythmias occurred in 2% of OHT patients leading to premature termination of the adenosine infusion.

OHT patients were excluded from the early trials of REG, which led to its approval, and thus the safety of REG in this population was initially unknown. The effects of REG in these patients are particularly relevant, however, given its relative $A_2A$ selectivity and the decreased incidence of AV block observed with REG in other populations. Cavalcante et al.[33] identified 40 OHT patients who underwent REG MPI. These results were compared with prior adenosine MPI results in the same patients. There were five episodes of second-degree AV block (Mobitz type II) and three episodes of sinus pause in adenosine MPI compared with only one episode of sinus pause in REG MPI. No major adverse effects such as congestive heart failure or death were reported following REG administration. To reverse REG’s side effects, aminophylline was given to four patients (two for severe headache and two for chest pressure). However, REG was largely well tolerated by the OHT patients with no difference in overall adverse effect profile between the two test drugs.

4. REG in positron emission tomography stress myocardial perfusion imaging

Although REG was approved in April 2008 by the U.S. Food and Drug Administration for use in single photon emission computed tomography (SPECT) radionuclide myocardial perfusion imaging (MPI) as a pharmacologic stressor in patients unable to perform exercise stress testing, it has not yet been formally approved for use in positron emission tomography (PET) MPI. Nonetheless, it is increasingly being used in PET MPI in addition to the more established vasodilators, adenosine and dipyridamole. Over the past several years, PET MPI has become more accepted into the mainstream for the diagnosis and management of coronary artery disease (CAD).[34] Furthermore, a recent consensus statement by the American Society of Nuclear Cardiology recommended PET MPI over SPECT MPI as the preferred initial pharmacologic MPI modality if available.[35] The following is a discussion of the current evidence for REG as a pharmacologic stressor in PET MPI.

4.1. Current guidelines

The 2003 ACC/AHA/ASNC Guidelines for Clinical Use of Radionuclide Imaging recommend adenosine or dipyridamole myocardial perfusion PET for diagnosis in patients with an intermediate likelihood of CAD and/or for risk stratification in patients with an intermediate or high likelihood of CAD.[36] The only class I recommendation is in “patients in whom an appropriately indicated myocardial perfusion SPECT study has been found to be equivocal for diagnostic or risk stratification purposes” (Level of Evidence B). Class IIa recommendations for vasodilator PET MPI are identification of “the extent, severity, and location of ischemia as the initial diagnostic test in patients who are unable to exercise” and in “patients who are able to exercise but have LBBB or an electronically-paced rhythm” (both Level of Evidence B). REG
is listed as an additional vasodilator in the 2009 American Society of Nuclear Cardiology Guidelines.[37]

4.2. PET vs. SPECT myocardial perfusion imaging: advantages and disadvantages

Cardiac PET imaging always includes concomitant CT acquisition for attenuation correction whereas this is still optional with SPECT. Effective radiation dose is lower with PET despite high positron emission energy due to very short half-life of rubidium-82 (Rb-82), the most commonly used PET radiotracer. Ejection fraction (EF) reserve (stress EF – rest EF) is more accurate with PET than SPECT because PET calculates the EF at peak stress rather than post stress as with SPECT. Coronary blood flow/flow reserve is possible with PET as myocardial uptake of Rb-82 bears a more linear relationship to coronary flow rates whereas the uptake of SPECT tracers plateaus at low flows. This allows for better characterization and localization of CAD. The superior image quality of PET is related to its high spatial resolution, reduced scatter, and the high positron emission energy of Rb-82 (1.52 MeV).

<table>
<thead>
<tr>
<th>Advantages</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>Higher spatial resolution (2–4 mm vs. 6–8 mm)</td>
<td>Incompatible with exercise (t1/2 of Rb-82 only 75 s)</td>
</tr>
<tr>
<td>Better count efficiency (more counts in less time)</td>
<td>Insurance coverage not universal</td>
</tr>
<tr>
<td>Superior soft tissue attenuation correction [38-39]</td>
<td>Less availability</td>
</tr>
<tr>
<td>Less liver/bowel uptake (less scatter) [40]</td>
<td>Motion artifact affects entire image (360° acquisition)</td>
</tr>
<tr>
<td>Shorter scan time (5 vs. 16 min) [40]</td>
<td>Claustrophobia (longer tunnel)</td>
</tr>
<tr>
<td>Less radiation (3.7 vs. 10–22 mSv) [40, 41]</td>
<td>More accurate estimation of EF reserve [42]</td>
</tr>
<tr>
<td>Ability to assess coronary blood flow/coronary flow</td>
<td>Superior diagnostic sensitivity, specificity, and accuracy [40]</td>
</tr>
<tr>
<td>Superior image quality [40]</td>
<td>Superior image quality [40]</td>
</tr>
<tr>
<td>Increased confidence in interpretation [40]</td>
<td></td>
</tr>
</tbody>
</table>

Table 2. Advantages and disadvantages of Rb-82 PET MPI vs. SPECT MPI

<table>
<thead>
<tr>
<th>Sensitivity (PET/SPECT)</th>
<th>Specificity (PET/SPECT)</th>
<th>Accuracy (PET/SPECT)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt;70% stenosis</td>
<td>87%/82% (rs)</td>
<td>93%/73% (P = 0.02)</td>
</tr>
<tr>
<td>&gt;50% stenosis</td>
<td>86%/81% (rs)</td>
<td>100%/66% (P = 0.00008)</td>
</tr>
</tbody>
</table>

Table 3. Overall sensitivity, specificity, and diagnostic accuracy of PET vs. SPECT MPI for both moderate and severe degrees of coronary stenosis

Comparison of 112 SPECT MPI (using adenosine and Tc-99m) and 112 PET MPI (using dipyridamole and Rb-82) in populations matched for gender, BMI, and presence/extent of
CAD.[40] “Specificity” includes low-likelihood patients who did not undergo angiography in addition to angiographically normal patients. “ns” = not statistically significant.

4.3. Advantages of REG in PET MPI

The increase in coronary blood flow is over 100 times greater with REG than adenosine. Rapid onset of hyperemia (less than 1 min after injection) with peak hyperemia occurring about 2.3 min following injection[3] along with weight-independent standardized dosing make REG well suited for use with short-acting PET radiotracers such as Rb-82 ($t_{1/2} = 75$ s). Rapid testing is thereby facilitated with the stress portion lasting less than 1 min. When using REG stress together with PET imaging, the entire test duration is only 16–18 min. Figure 3 is a flow diagram of the REG PET MPI protocol used by Hsiao et al.[43]


Figure 3. Rest-stress regadenoson [82]Rb PET/CT protocol. After scout CT acquisition (120 kVp, 10 mA), CT transmission scan (CTAC) (140 kVp, 10 mA, pitch of 1.35) was acquired. Patients received 1,480-2,220 MBq of [82]Rb intravenously at rest, and emission images were acquired in 2-dimensional list mode. After rest imaging, patients remained in scanner gantry for stress imaging. Stress was induced with 0.4 mcg of regadenoson given intravenously over 10 s followed by 10-mL flush with normal saline. Immediately after saline flush, second dose of 1,480-2,220 MBq of [82]Rb was administered intravenously approximately 30 s after regadenoson injection and emission images were acquired as previously described. Ordered-subsets expectation maximization (30 iterations and 2 subsets) and 3-dimensional PET filtering (Butterworth filter, cutoff frequency of 10, order of 5) were used for reconstruction of images.[43]

4.4. Coronary flow reserve using PET

The ability to quantitatively assess coronary blood flow (CBF) and coronary flow reserve (CFR) on angiography was discovered by Gould in animal experiments during the mid 1970s.[44] Because PET image acquisition occurs during peak stress, calculation of CFR (peak flow ÷ rest flow) is one of the unique features of PET as opposed to other noninvasive imaging modalities. A “normal range” has proved difficult to define given the disparity between coronary flows in asymptomatic patients. Based on pooled data from nearly 15,000 patients in 252 studies using three different PET isotopes, CFR in patients without CAD is 3.55 ± 1.36. In patients with established coronary disease, this drops to 2.02 ± 0.70.[45] Table 4 displays the range of values...
for absolute coronary flow and CFR in the presence of CAD risk factors and other forms of cardiac disease. One of the larger studies in the literature, however, identified a CFR of 1.74 as the cutoff for “definite ischemia” below which patients manifest anginal symptoms and/or ischemic ECG changes during vasodilator stress testing matched by a significant perfusion defect on PET imaging.[46]


Table 4. Graded Absolute Flow and Coronary Flow Reserve Across Spectrum of Disease: n=14,962[45]

<table>
<thead>
<tr>
<th>Population</th>
<th>n</th>
<th>Rest Flow (cc/min/g)</th>
<th>Stress Flow (cc/min/g)</th>
<th>CFR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal controls</td>
<td>3,484</td>
<td>0.82 ± 0.06</td>
<td>2.86 ± 1.29</td>
<td>3.55 ± 1.36</td>
</tr>
<tr>
<td>Risk factors only</td>
<td>3,592</td>
<td>0.85 ± 0.08</td>
<td>2.25 ± 1.07</td>
<td>2.80 ± 1.39</td>
</tr>
<tr>
<td>Established coronary artery disease</td>
<td>1,650</td>
<td>0.83 ± 0.10</td>
<td>1.71 ± 0.71</td>
<td>2.02 ± 0.70</td>
</tr>
<tr>
<td>Mixed (risk factors and/or known coronary artery disease)</td>
<td>4,765</td>
<td>0.97 ± 0.10</td>
<td>1.86 ± 0.58</td>
<td>1.93 ± 0.48</td>
</tr>
<tr>
<td>Cardiomyopathy</td>
<td>594</td>
<td>0.73 ± 0.07</td>
<td>1.47 ± 0.56</td>
<td>2.02 ± 0.67</td>
</tr>
<tr>
<td>Hypertrophic cardiomyopathy</td>
<td>345</td>
<td>0.90 ± 0.10</td>
<td>1.57 ± 0.33</td>
<td>1.84 ± 0.38</td>
</tr>
<tr>
<td>Syndrome X</td>
<td>348</td>
<td>1.06 ± 0.11</td>
<td>2.65 ± 1.31</td>
<td>2.54 ± 1.31</td>
</tr>
<tr>
<td>After cardiac transplant</td>
<td>184</td>
<td>1.14 ± 0.18</td>
<td>2.44 ± 1.34</td>
<td>2.29 ± 0.86</td>
</tr>
</tbody>
</table>

N = 14,962 from 252 unique publications. N:13 ammonia = 5,541; 0:15 water = 3,161; Rb-82 = 6,171.


Table 4. Graded Absolute Flow and Coronary Flow Reserve Across Spectrum of Disease: n=14,962[45]

Not until recently was the clinical utility of PET-derived CFR fully appreciated. A study of 205 patients (using REG in half of these) demonstrated that with a negative predictive value of 97%, normal global CFR virtually assures the absence of high-risk CAD, despite any coexistent abnormal perfusion.[47] However, as a reduced CFR can occur in three different conditions (diffuse non-obstructive atherosclerosis, significant epicardial coronary stenosis, and microvascular disease), it can be somewhat helpful but is not specific for selecting patients likely to have high-risk CAD on angiography. A very recent study of PET-derived CFR further illustrated that low CFR can be seen in patients with systolic cardiomyopathy (EF ≤ 45%) of both ischemic and non-ischemic etiologies.[48] Murthy et al.[49] studied 2783 patients with known or suspected CAD referred for rest/stress PET MPI and then followed over a median of 1.4 years. Those in the lowest CFR tertile (<1.5) had a 16-fold increase in risk of cardiac death versus those in the highest tertile (>2.0). The middle tertile had a 5.7-fold increase in risk compared to the highest tertile. The addition of CFR to clinical and standard MPI factors led to the correct re-categorization of 34.9% of patients in the intermediate-risk group. Patients in this study received one of four different vasodilators (adenosine, dipyridamole, dobutamine, or REG). As resting CBF was similar between all three tertiles, the reduction in CFR was primarily driven by lower CBF with stress suggesting impaired coronary vasodilator function as an etiology. No difference was drawn between the various vasodilators used, however.

Very little has been published on the specific use of REG to assess CFR in Rb-82 PET MPI. Van Tosh et al.[50] used REG alone to show that CFR corresponded with LV dysfunction (LVD) during stress and that regional reductions in CFR were more often present in patients with
LVD than those without, indicating that the phenomenon of coronary steal may be involved in the genesis of LVD.

4.5. REG PET MPI vs. dipyridamole PET MPI

There exist scant data comparing REG and dipyridamole in PET MPI. A recent study retrospectively assessed CBF and CFR using Rb-82 perfusion PET/CT in 104 matched patients with normal stress tests, half with dipyridamole and half with REG. No significant difference in stress CBF and CFR was found between the two vasodilators (Figure 4). Further supporting REG’s usefulness as a stress agent was the lack of any correlation between stress CBF or CFR and patient weight or BMI.[51]

A very recent study by Johnson and Gould compared CFR in patients undergoing two sequential PET MPIs, either both with dipyridamole (n = 50) or with dipyridamole and REG


Figure 4. Myocardial blood flow (MBF) and myocardial flow reserve (MFR) in subjects undergoing pharmacological stress with regadenoson versus dipyridamole. a No significant difference in MBF between groups at rest (left, p=0.77) or during stress (right, p=0.39). b No significant difference in MFR (p=0.31).[51]
In the latter group, various timings between REG administration and activation of the Rb-82 generator were used. It was demonstrated that using the timing recommended in the REG package insert (10–20 s between REG injection and radioisotope injection), the stress CBF and CFR with REG were only 80% of the hyperemia attained with dipyridamole. By increasing this interval to 55 ms, this percentage was increased to 90%. These findings suggest that with the current timing recommendation, REG remains inferior to dipyridamole in detecting stress CBF and CFR.

The shorter duration of peak hyperemia with REG (2.3 min) than dipyridamole has raised some concern as to whether Rb-82 uptake by the myocardium would be sufficient to register perfusion defects or changes in cardiac function with the newer vasodilator. Cullom et al.[53] studied 32 patients, all of whom underwent both REG and dipyridamole PET MPI, and compared summed stress and difference scores, total perfusion deficit, LVEF, LV volumes, and change in stress-rest function. They determined that REG and dipyridamole yielded equivalent measures of cardiac perfusion and function.

To date, there are no published investigations of REG vs. adenosine in PET myocardial perfusion imaging.

4.6. Diagnostic accuracy of REG PET MPI

Studies comparing vasodilator stress SPECT and PET MPI have repeatedly demonstrated slightly higher sensitivity in PET (90%) than SPECT (80–84%) but far greater specificity in PET (89%) than SPECT (53–76%).[34, 38, 39] Table 5 summarizes the results of all published literature on the diagnostic accuracy of PET through 2007. Most of these studies used Rb-82 as a tracer and dipyridamole ± handgrip for stress. One included dipyridamole, adenosine, and dobutamine stress, and one used exercise stress with ammonia-N13 PET imaging.

<table>
<thead>
<tr>
<th>Reference</th>
<th>No. of patients</th>
<th>Women</th>
<th>Prior CAD</th>
<th>PET radiotracer</th>
<th>Sensitivity</th>
<th>Specificity</th>
<th>PPV</th>
<th>NPV</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sampson et al. (23)</td>
<td>102</td>
<td>0.42</td>
<td>0</td>
<td>124 Rb</td>
<td>0.93</td>
<td>0.83</td>
<td>0.90</td>
<td>0.94</td>
<td>0.87</td>
</tr>
<tr>
<td>Bateman et al. (21)</td>
<td>112</td>
<td>0.46</td>
<td>0.25</td>
<td>124 Rb</td>
<td>0.87</td>
<td>0.93</td>
<td>0.95</td>
<td>0.84</td>
<td>0.89</td>
</tr>
<tr>
<td>Marwick et al. (23)</td>
<td>74</td>
<td>0.19</td>
<td>0.49</td>
<td>124 Rb</td>
<td>0.90</td>
<td>1</td>
<td>1</td>
<td>0.36</td>
<td>0.91</td>
</tr>
<tr>
<td>Grover-McKay et al. (24)</td>
<td>31</td>
<td>0.01</td>
<td>0.13</td>
<td>124 Rb</td>
<td>1</td>
<td>0.73</td>
<td>0.80</td>
<td>1</td>
<td>0.87</td>
</tr>
<tr>
<td>Stewart et al. (25)</td>
<td>81</td>
<td>0.36</td>
<td>0.42</td>
<td>124 Rb</td>
<td>0.83</td>
<td>0.86</td>
<td>0.94</td>
<td>0.64</td>
<td>0.84</td>
</tr>
<tr>
<td>Go et al. (19)</td>
<td>202</td>
<td>0.47</td>
<td>0</td>
<td>124 Rb</td>
<td>0.93</td>
<td>0.78</td>
<td>0.93</td>
<td>0.80</td>
<td>0.90</td>
</tr>
<tr>
<td>Demer et al. (25)</td>
<td>193</td>
<td>0.26</td>
<td>0.34</td>
<td>124 Rb /15O-ammonia</td>
<td>0.83</td>
<td>0.95</td>
<td>0.96</td>
<td>0.90</td>
<td>0.86</td>
</tr>
<tr>
<td>Tamaki et al. (26)</td>
<td>51</td>
<td>0.75</td>
<td>0</td>
<td>124 Rb /15O-ammonia</td>
<td>0.98</td>
<td>1</td>
<td>1</td>
<td>0.75</td>
<td>0.98</td>
</tr>
<tr>
<td>Gould et al. (27)</td>
<td>31</td>
<td>0.98</td>
<td>0</td>
<td>124 Rb /15O-ammonia</td>
<td>0.95</td>
<td>1</td>
<td>1</td>
<td>0.90</td>
<td>0.97</td>
</tr>
<tr>
<td>Weighted summary</td>
<td>877</td>
<td>0.29</td>
<td>0.35</td>
<td>124 Rb /15O-ammonia</td>
<td>0.90</td>
<td>0.89</td>
<td>0.94</td>
<td>0.73</td>
<td>0.90</td>
</tr>
</tbody>
</table>

*Study using PET/CT (in which CT was used for attenuation correction only).

PPV= positive predictive value; NPV= negative predictive value; NR= not reported. (Reprinted with permission of (28).)
Hsiao et al. [43] performed the first and so far only published study to evaluate the diagnostic accuracy of REG in PET MPI. In a relatively small cohort of 134 patients in 98 of whom angiographic data were also available, its accuracy was found to be similar to that of PET MPI using other vasodilators. Sensitivity for obstructive CAD was 92%, and overall specificity was 77% (53% in patients with high likelihood of CAD but no angiographic evidence of obstructive disease and 93% in low likelihood patients who did not go on to angiography [normalcy rate]). The area under the receiver–operator curve was 0.847, comparable to the high accuracy rates of PET in previous studies.

The high sensitivity of PET MPI for detection of obstructive CAD can be further increased by PET’s ability to quantify blood flow/flow reserve and to calculate LVEF reserve using peak-stress LVEF.

4.7. Prognostic value of REG PET MPI

It has been shown that the prognostic value of REG is comparable to that of adenosine in patients with normal SPECT myocardial perfusion tests.[54] There are no published data on the prognostic value of REG in PET MPI, nor of REG MPI in patients with abnormal results using either PET or SPECT. Recent studies, however, offer insight as to the prognostic value of LVEF reserve in vasodilator stress PET.

Dorbala et al. [42] established that LVEF reserve (stress LVEF – rest LVEF) is independently predictive of the extent of at-risk myocardium on Rb-82 PET MPI and the extent of CAD on invasive angiography. Based on these results, LVEF reserve >5% essentially rules out severe 3-vessel or left main disease with a negative predictive value of 97%. In 985 patients with gated vasodilator stress Rb-82 PET MPI, nearly half of whom were at intermediate risk for CAD consistent with contemporary practice, the same group of investigators showed that during a mean follow-up period of 1.7 years, the frequency of cardiac events and all-cause death was higher in patients with LVEF reserve <0 than in those with LVEF which either remained the same or augmented with stress.[55] The prognostic value of LVEF reserve was found to be independent of, and incremental to, clinical variables and rest LVEF. These studies, however, included only patients who had received either dipyridamole or adenosine.

Hsiao’s was the first group to investigate LVEF reserve using REG PET MPI, albeit in a much smaller cohort of 115 patients. Here, LVEF reserve with REG was inversely related to the severity of reversible perfusion defect (summed difference score) as well as jeopardized myocardium on coronary angiography (Duke Jeopardy Score)[43] (Figures 5 and 6). This suggests that REG may be as useful as dipyridamole or adenosine in determining LVEF reserve; however, further studies are still needed to evaluate its prognostic value.

4.8. Future directions for REG PET MPI

The IDEALPET (Integrated Dual Exercise and Lexiscan PET) study is currently underway and will compare Lexiscan© alone with Lexiscan© plus exercise (“Lexercise”) with regards to safety, tolerability, myocardial perfusion image quality, and assessment of relative and absolute myocardial perfusion.[56]

Figure 5. Regadenoson LVEF reserve as function of relative MPI results. Mod=moderate.[43]


Figure 6. Regadenoson LVEF reserve as function of Duke Jeopardy Score. LLK= low likelihood.[43]
As of February 2011 REG was being used in 68% of all pharmacologic stress MPI studies in the United States.[54] Given its already widespread use and favorable profile as a stress agent plus the advantages inherent in Rb-82 PET perfusion imaging (superior image quality, shorter scan time, lower radiation dose to patient, quantitation of myocardial blood flow, measurement of peak LVEF, additional prognostic information), REG PET MPI has the capacity to become the pharmacologic stress test of choice over the next several years.

5. Novel applications of REG

5.1. Adjunct to exercise MPI

Exercise-based testing has been convincingly shown to provide powerful prognostic data and remains the preferred mode of stress testing if patients are capable of exercising.[57, 58] However, about 25% of exercise-based testing may be non-diagnostic due to inability to achieve target heart rates. Two alternatives for these patients have been evaluated in the past: either rescheduling for pharmacologic stress or immediately attempting adjunctive vasodilator stress with agents such as adenosine and dipyridamole.[59, 60] The combination of simultaneous adjunctive low-level exercise with adenosine or dipyridamole helps both to lessen side effects and improve image quality.[61, 62] However, trying to add on adenosine or dipyridamole when exercise testing is submaximal poses major challenges as both are given as an infusion over a few minutes, need to be adjusted for weight or delivered via pump (as in the case of adenosine), and thus are not immediately feasible. In these instances, patients are usually rescheduled for a pharmacologic stress test when exercise testing is submaximal.

With the advent of rapid-acting, weight-independent, single-bolus dosing of REG, its use as an adjunct to exercise seemed logistically feasible and potentially convenient. Its administration could result in quick conversion of an otherwise non-diagnostic nuclear exercise stress study due to submaximal heart rate to a diagnostic one. Early data support such a practice.

Thomas et al.[63] evaluated the safety of REG during exercise in a double blind study of 60 patients focusing on image quality, patient acceptance, and detection of perfusion defects. Patients undergoing a clinically indicated adenosine supine MPI were subsequently randomized in a 2:1 fashion to REG with low-level exercise (RegEx) or placebo with low-level exercise (PlcEx). This small study showed no significant differences in blood pressure response between the RegEx and PlcEx groups, although a smaller increase in heart rate was noted in the RegEx than in the PlcEx group. The image quality was better with RegEx compared to the adenosine supine MPI images. Patient tolerability was also reported to be better with RegEx compared to adenosine supine MPI. No significant adverse events, including high-grade AV block, were reported in the RegEx group.

In a subsequent study, Kwon et al.[27] published their retrospective experience with 1263 patients undergoing REG MPI with either adjunctive low-level treadmill exercise (n = 596) or as a standard supine REG stress test (n = 667). Among all participants an asymptomatic drop in systolic blood pressure > 10 mmHg occurred in 51% and > 30 mmHg in 9%. A pressure drop was observed more often in those randomized to REG plus low-level treadmill exercise (56%) than in those undergoing supine REG (47%). In their COPD/asthma patients
(16%), REG with low-level exercise was well tolerated, and they also reported lower incidence of nausea, shortness of breath, transient heart block, palpitations, and dizziness overall in those who underwent low-level exercise.

Our own experience comparing REG MPI \((n = 887)\) to REGWALK MPI \((n = 485)\) (REG with adjunctive low-level exercise) was published as a retrospective series. We showed that REGWALK studies demonstrated higher stress heart rate response, higher heart rate reserve, and higher systolic blood pressure with stress. There was less use of aminophylline for reversal of REG side effects in the REGWALK compared to the REG group. No major adverse events were reported in this series.

No data exist showing improved detection of ischemia/prognosis by combining REG with exercise. A few randomized studies have assessed the safety and efficacy of REG when used as an adjunct to maximal exercise when target heart rate is not achieved. Ross et al.[64] randomized 200 patients undergoing exercise MPI to either adjunctive REG if target heart rate was not achieved at peak exercise or to the discontinuation of exercise with conversion to a standard supine REG stress test. They showed that both approaches were well tolerated without any adverse events. There were no differences in ischemia detection, image quality or referral to cardiac catheterization in either group. Another small randomized study \((n = 140)\) also showed that augmenting submaximal exercise with REG as needed was safe in patients.[65] In an effort to finalize the evaluation of REG’s safety as an adjunct to exercise, a large randomized trial has just been completed by Astellas.[66] Results of this study will conclusively address not only the safety of REG with exercise but also the detection of ischemia when compared to REG alone.

5.2. Fractional Flow Reserve (FFR)

The concept of reactive hyperemia is particularly useful in guiding percutaneous coronary intervention (PCI) when intermediate coronary lesions of unclear hemodynamic significance are present on invasive angiography.[67, 68] More recently, seminal studies have firmly established that FFR-guided decision making for coronary lesions of unclear significance is associated with a favorable outcome with PCI being deferred or performed based on FFR values.[69] Most catheterization labs use either intracoronary or intravenous adenosine for assessment of hyperemic response.[70, 71] However recent studies have now shown that REG may be a viable alternative to adenosine with its weight independent bolus and rapid achievement of hyperemia in 33–40 s, thus shortening the entire time needed for FFR assessment.[72, 73] In a study of 25 patients undergoing catheterization, Nair et al.[72] compared the ability of IV adenosine and IV REG to induce coronary hyperemia in assessment of coronary stenosis significance. They found excellent linear correlation for measurement of FFR between the two agents \((r = 0.985, P = 0.001)\). Furthermore, none of the hemodynamically significant lesions (FFR <0.8, 52% of patients) identified by adenosine were reclassified by REG. There were no significant adverse reactions to either drug and REG was overall better tolerated than adenosine.[72] In a more recent study by Prasad et al.,[74] the authors compared 57 patients (60 lesions) undergoing FFR measurements first with adenosine followed by a 10 min washout phase and then with REG. They showed high correlation in hyperemic response between the two drugs \((R^2 = 0.93)\) (Figure 7) and substantially shorter time to peak hyperemia with REG
than adenosine as well as a trend to a better side effect profile with REG. One issue of concern raised by these authors has been the potential cost of a single vial of REG (around 250 dollars) compared to a 3-min adenosine infusion (80 dollars). However, such cost differences could be made up by shorter duration of REG administration, no need for infusion pumps and less nursing time for set up. A recent randomized study of 100 patients has also shown that REG is equivalent to central venous infusion of adenosine to induce maximal hyperemia for FFR determination.[75]

In summary, the data accumulated on REG in FFR suggest that it could very well be the preferred agent in the catheterization lab given its ease of use and proven efficacy and comparability to adenosine.

5.3. Stress echocardiography

The detection of CAD using stress echocardiography (SE) is based on the physiologic principle of stress-induced subendocardial ischemia causing wall motion abnormalities in the territory subtended by stenosis. Exercise and dobutamine (DSE) are the main methods of SE in North America,[76] whereas high-dose dipyridamole supplemented with atropine has been the mainstay pharmacologic stressor in Europe.[76, 77]

It is well known that wall motion can be completely normal with DSE despite mild to moderate stenosis and corresponding abnormalities in hyperemic blood flow.[78] Using a newer technique of myocardial contrast echocardiography (MCE), contrast imaging during induced hyperemia allows for the detection of milder degrees of coronary stenosis. Similar to nuclear perfusion imaging, MCE is able to pick up small perfusion abnormalities, which occur prior to ischemic changes in wall motion, in keeping with the “ischemic cascade.” Prior studies using adenosine, dobutamine, and exercise with MCE have shown that myocardial contrast perfusion enables detection of moderate stenosis when added to wall motion.[79–81]

Given its ease of use, there has been interest in using REG as a vasodilator to induce hyperemic stress during MCE. In a study of 100 patients undergoing quantitative coronary angiography, Porter et al.[82] performed real-time MCE with Definity 3% infusion at baseline and then at 2-min intervals for up to 6 min after a REG bolus. This study showed that MCE with REG can detect noncritical coronary stenosis (>50% diameter) with sensitivity, specificity, and accuracy of 80%, 74%, and 75%, respectively, which was better than wall motion analysis alone (60%, 70%, and 66%, respectively (P < 0.001 for sensitivity)). Furthermore, the authors concluded that the sensitivity was highest when imaging was performed 4–6 min after REG administration.[82] In a recent study performed in 10 dogs with mild to moderate non-flow limiting CAD, Le et al.[83] used REG (5 μg/kg, 10-s bolus) along with MCE and assessed myocardial blood volume, flow velocity, and total regional myocardial flow before and after REG administration. REG induced an increase in coronary blood flow for 30 min. This decreased proportionally to stenosis severity, and perfusion defects were visible for up to 10 min after REG bolus. They noted that the optimal time for imaging myocardial perfusion in stress echo with REG was between 3 and 10 min after REG bolus.[83]
Our group recently reported on 44 patients undergoing diagnostic angiography based on prior abnormal stress testing who also underwent a novel protocol called REGAT (REG + atropine) SE to assess feasibility, safety, and diagnostic accuracy of CAD detection. The testing sequence began with administration of $2 \times 1$ mg boluses of atropine to induce chronotrope.

Figure 7. Average of FFR with Adenosine and FFR with Regadenoson[74]
py followed by a 400-μg bolus of REG, and then echo imaging at peak stress starting 20 s after the REG bolus. The protocol was found to be safe and well tolerated with no serious adverse effects. The mean duration of REGAT SE was 18 ± 7 min. Significant CAD (≥70% stenosis) by angiography was present in 51.1%. Sensitivity, specificity, and positive and negative predictive values for REGAT SE were 60.9%, 80.4%, 82.4%, and 67.9%, respectively. By coronary territories, the sensitivity, specificity, PPV, and NPV were as follows: left anterior descending artery, 58.8%, 92.9%, 83.3%, and 78.8%; left circumflex artery, 6.7%, 93.3%, 33.3%, and 67.7%; and right coronary artery, 16.7%, 93.9%, 50%, and 75.6%. Over 90% of subjects reported feeling comfortable, with 83% preferring REGAT as a future stress modality. We concluded that although the REGAT protocol was fast, safe, and well-tolerated, with good specificity for CAD detection, its low sensitivity and NPV preclude it from routine use. Importantly, contrast was not utilized in our study as we were testing the feasibility of a combination of REG and atropine. Overall evidence indicates that REG in SE may be feasible and safe and, but larger studies are needed in this area as concern still exits that echocardiographic imaging may not detect ischemia induced by vasodilator stress.

5.4. Coronary CT Angiography (CCTA) and stress perfusion

It is now well established that CCTA performs with high diagnostic sensitivity and has excellent negative predictive value for the noninvasive evaluation of CAD[84, 85] However the specificity and positive predictive value have been shown to be less than desired with overestimation of stenosis severity in published studies[86, 87]. When compared to fractional flow reserve or SPECT even apparent high grade stenosis diagnosed on CCTA has not be consistently associated with ischemia[87]. This has raised some concerns that CCTA as a noninvasive modality for CAD may lead to higher false positives and downstream testing. CCTA stenosis detection requires additional physiologic information to correctly identify physiologic significant lesions. Until recently evidence for ischemia evaluation with CCTA has been very limited. The concept of combining stress perfusion with CT (CTP) has been tested and found to be accomplished in many single center studies mainly using adenosine or adenosine triphosphate. This has raised the possibility that a comprehensive anatomic and physiologically CAD assessment could be feasible by CCTA+CTP. [88-90]

Most recently a multicenter study sponsored by Astellas was completed and published evaluating the non-inferiority of REG CTP to REG SPECT. Patients (men > 45 years; women > 50 years) with known or suspected coronary artery disease (n=124) were randomized to 1 of 2 diagnostic sequences: rest/REG SPECT MPI on day 1, then REG/rest CTP on day 2, or REG/rest CTP on day 1 followed by rest/REG SPECT MPI on day 2. CCTA was also performed during the same acquisition as the CTP in both groups. Scanning platforms included 64-, 128-, 256-, and 320-slice systems. The primary analysis examined the agreement rate between CTP and SPECT for detecting or excluding reversible ischemia in 2 myocardial segments as assessed by independent blinded readers. Across the 110 patients included in the final analysis REG CTP was non-inferior to SPECT for detecting or excluding reversible ischemia with an agreement rate of 0.87 (95% confidence interval [CI], 0.77-0.97) and sensitivity and specificity of 0.90 (95% CI, 0.71-1.00) and 0.84 (95% CI, 0.77-0.91), respectively. The agreement rate for
detecting or excluding fixed defects by REG CTP and SPECT was 0.86 (95% CI, 0.74-0.98). With SPECT as the reference standard, the diagnostic accuracies for detecting or excluding ischemia by REG CTP and CTA alone were 0.85 (95% CI, 0.78-0.91) and 0.69 (95% CI, 0.60-0.77), respectively. The authors concluded that REG CTP is non-inferior to SPECT. Thus, CT vasodilator stress perfusion imaging either with REG or adenosine appears to have a promising role in providing physiologic information to clarify anatomic stenosis. Further studies are awaited to establish this modality in clinical practice.

6. Conclusion

We have aimed to provide the reader in this chapter a detailed overview of REG and its current status in cardiac stress testing and other emerging cardiac applications. The role of REG remains to be better defined in cardiac MRI and CT.

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