1 ADENOSCAN®

2 (adenosine injection)

3 FOR INTRAVENOUS INFUSION ONLY

4 Revised: April 2005

5 **DESCRIPTION**

- 6 Adenosine is an endogenous nucleoside occurring in all cells of the body. It is chemically 6-
- 7 amino-9-beta-D-ribofuranosyl-9-H-purine and has the following structural formula:



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C₁₀H₁₃N₅O₄ 267.24

10 Adenosine is a white crystalline powder. It is soluble in water and practically insoluble in

11 alcohol. Solubility increases by warming and lowering the pH of the solution.

Each Adenoscan vial contains a sterile, non-pyrogenic solution of adenosine 3 mg/mL and
sodium chloride 9 mg/mL in Water for Injection, q.s. The pH of the solution is between 4.5 and

14 7.5.

15 CLINICAL PHARMACOLOGY

16 Mechanism of Action

17 Adenosine is a potent vasodilator in most vascular beds, except in renal afferent arterioles and 18 hepatic veins where it produces vasoconstriction. Adenosine is thought to exert its pharmacological effects through activation of purine receptors (cell-surface A1 and A2 adenosine 19 20 receptors). Although the exact mechanism by which adenosine receptor activation relaxes 21 vascular smooth muscle is not known, there is evidence to support both inhibition of the slow 22 inward calcium current reducing calcium uptake, and activation of adenylate cyclase through A₂ 23 receptors in smooth muscle cells. Adenosine may also lessen vascular tone by modulating 24 sympathetic neurotransmission. The intracellular uptake of adenosine is mediated by a specific 25 transmembrane nucleoside transport system. Once inside the cell, adenosine is rapidly 26 phosphorylated by adenosine kinase to adenosine monophosphate, or deaminated by adenosine 27 deaminase to inosine. These intracellular metabolites of adenosine are not vasoactive. 28 Myocardial uptake of thallium-201 is directly proportional to coronary blood flow. Since 29 Adenoscan significantly increases blood flow in normal coronary arteries with little or no 30 increase in stenotic arteries, Adenoscan causes relatively less thallium-201 uptake in vascular

31 territories supplied by stenotic coronary arteries i.e., a greater difference is seen after Adenoscan

32 between areas served by normal and areas served by stenotic vessels than is seen prior to

33 Adenoscan.

34 Hemodynamics

Adenosine produces a direct negative chronotropic, dromotropic and inotropic effect on the
 heart, presumably due to A₁-receptor agonism, and produces peripheral vasodilation, presumably

due to A₂-receptor agonism. The net effect of Adenoscan in humans is typically a mild to
 moderate reduction in systolic, diastolic and mean arterial blood pressure associated with a reflex
 increase in heart rate. Rarely, significant hypotension and tachycardia have been observed.

40 **Pharmacokinetics**

41 Intravenously administered adenosine is rapidly cleared from the circulation via cellular uptake, 42 primarily by erythrocytes and vascular endothelial cells. This process involves a specific 43 transmembrane nucleoside carrier system that is irreversible, nonconcentrative, and 44 bidirectionally symmetrical. Intracellular adenosine is rapidly metabolized either via 45 phosphorylation to adenosine monophosphate by adenosine kinase, or via deamination to inosine by adenosine deaminase in the cytosol. Since adenosine kinase has a lower K_m and V_{max} than 46 47 adenosine deaminase, deamination plays a significant role only when cytosolic adenosine 48 saturates the phosphorylation pathway. Inosine formed by deamination of adenosine can leave 49 the cell intact or can be degraded to hypoxanthine, xanthine, and ultimately uric acid. Adenosine 50 monophosphate formed by phosphorylation of adenosine is incorporated into the high-energy 51 phosphate pool. While extracellular adenosine is primarily cleared by cellular uptake with a half-52 life of less than 10 seconds in whole blood, excessive amounts may be deaminated by an ecto-53 form of adenosine deaminase. As Adenoscan requires no hepatic or renal function for its 54 activation or inactivation, hepatic and renal failure would not be expected to alter its 55 effectiveness or tolerability.

56 **Clinical Trials**

In two crossover comparative studies involving 319 subjects who could exercise (including 106
healthy volunteers and 213 patients with known or suspected coronary disease), Adenoscan and

59 exercise thallium images were compared by blinded observers. The images were concordant for 60 the presence of perfusion defects in 85.5% of cases by global analysis (patient by patient) and up 61 to 93% of cases based on vascular territories. In these two studies, 193 patients also had recent 62 coronary arteriography for comparison (healthy volunteers were not catheterized). The 63 sensitivity (true positive Adenoscan divided by the number of patients with positive (abnormal) 64 angiography) for detecting angiographically significant disease (>50% reduction in the luminal 65 diameter of at least one more vessel) was 64% for Adenoscan and 64% for exercise testing, while the specificity (true negative divided by the number of patients with negative angiograms) 66 67 was 54% for Adenoscan and 65% for exercise testing. The 95% confidence limits for Adenoscan 68 sensitivity were 56% to 78% and for specificity were 37% to 71%.

69 Intracoronary doppler flow catheter studies have demonstrated that a dose of intravenous

70 Adenoscan of 140 mcg/kg/min produces maximum coronary hyperemia (relative to

71 intracoronary papaverine) in approximately 95% of cases within two to three minutes of the

72 onset of infusion. Coronary blood flow velocity returns to basal levels within one to two minutes

73 of discontinuing the Adenoscan infusion.

74 INDICATIONS AND USAGE

75 Intravenous Adenoscan is indicated as an adjunct to thallium-201 myocardial perfusion

respective to exercise adequately (see Warnings).

77 CONTRAINDICTIONS

78 Intravenous Adenoscan (adenosine injection) should not be administered to individuals with:

79	1.	Second- or third-degree AV block (except in patients with a functioning artificial
80		pacemaker).

81 2. Sinus node disease, such as sick sinus syndrome or symptomatic bradycardia (except in
82 patients with a functioning artificial pacemaker).

- 83 3. Known or suspected bronchoconstrictive or bronchospastic lung disease (e.g., asthma).
- 84 4. Known hypersensitivity to adenosine.

85 WARNINGS

86 Fatal Cardiac Arrest, Life Threatening Ventricular Arrhythmias, and Myocardial

87 Infarction.

Fatal cardiac arrest, sustained ventricular tachycardia (requiring resuscitation), and nonfatal
myocardial infarction have been reported coincident with Adenoscan infusion. Patients with
unstable angina may be at greater risk. Appropriate resuscitative measures should be available.

91 Sinoatrial and Atrioventricular Nodal Block

92 Adenoscan (adenosine injection) exerts a direct depressant effect on the SA and AV nodes and 93 has the potential to cause first-, second- or third-degree AV block, or sinus bradycardia. 94 Approximately 6.3% of patients develop AV block with Adenoscan, including first-degree 95 (2.9%), second-degree (2.6%), and third-degree (0.8%) heart block. All episodes of AV block 96 have been asymptomatic, transient, and did not require intervention. Adenoscan can cause sinus 97 bradycardia. Adenoscan should be used with caution in patients with pre-existing first-degree 98 AV block or bundle branch block and should be avoided in patients with high-grade AV block or 99 sinus node dysfunction (except in patients with a functioning artificial pacemaker). Adenoscan

- 100 should be discontinued in any patient who develops persistent or symptomatic high-grade AV
- 101 block. Sinus pause has been rarely observed with adenosine infusions.

102 Hypotension

103 Adenoscan (adenosine injection) is a potent peripheral vasodilator and can cause significant

- 104 hypotension. Patients with an intact baroreceptor reflex mechanism are able to maintain blood
- 105 pressure and tissue perfusion in response to Adenoscan by increasing heart rate and cardiac
- 106 output. However, Adenoscan should be used with caution in patients with autonomic
- 107 dysfunction, stenotic valvular heart disease, pericarditis or pericardial effusions, stenotic carotid
- 108 artery disease with cerebrovascular insufficiency, or uncorrected hypovolemia, due to the risk of
- 109 hypotensive complications in these patients. Adenoscan should be discontinued in any patient
- 110 who develops persistent or symptomatic hypotension.

111 Hypertension

Increases in systolic and diastolic pressure have been observed (as great as 140 mm Hg systolic in one case) concomitant with Adenoscan infusion; most increases resolved spontaneously within several minutes, but in some cases, hypertension lasted for several hours.

115 Bronchoconstriction

116 Adenoscan (adenosine injection) is a respiratory stimulant (probably through activation of

- 117 carotid body chemoreceptors) and intravenous administration in man has been shown to increase
- 118 minute ventilation (Ve) and reduce arterial PCO₂ causing respiratory alkalosis. Approximately
- 119 28% of patients experience breathlessness (dyspnea) or an urge to breathe deeply with
- 120 Adenoscan. These respiratory complaints are transient and only rarely require intervention.

Adenosine administered by inhalation has been reported to cause bronchoconstriction in 121 122 asthmatic patients, presumably due to mast cell degranulation and histamine release. These 123 effects have not been observed in normal subjects. Adenoscan has been administered to a limited 124 number of patients with asthma and mild to moderate exacerbation of their symptoms has been 125 reported. Respiratory compromise has occurred during adenosine infusion in patients with 126 obstructive pulmonary disease. Adenoscan should be used with caution in patients with 127 obstructive lung disease not associated with bronchoconstriction (e.g., emphysema, bronchitis, 128 etc.) and should be avoided in patients with bronchoconstriction and bronchospasm (e.g. 129 asthma). Adenoscan should be discontinued in any patient who develops severe respiratory 130 difficulties.

131 **PRECAUTIONS**

132 **Drug Interactions**

Intravenous Adenoscan (adenosine injection) has been given with other cardioactive drugs (such as beta adrenergic blocking agents, cardiac glycosides, and calcium channel blockers) without apparent adverse interactions, but its effectiveness with these agents has not been systematically evaluated. Because of the potential for additive or synergistic depressant effects on the SA and AV nodes, however, Adenoscan should be used with caution in the presence of these agents.

The vasoactive effects of Adenoscan are inhibited by adenosine receptor antagonists, such as methylxanthines (e.g., caffeine and theophylline). The safety and efficacy of Adenoscan in the presence of these agents has not been systematically evaluated.

- 141 The vasoactive effects of Adenoscan are potentiated by nucleoside transport inhibitors, such as
- 142 dipyridamole. The safety and efficacy of Adenoscan in the presence of dipyridamole has not
- 143 been systematically evaluated.
- 144 Whenever possible, drugs that might inhibit or augment the effects of adenosine should be
- 145 withheld for at least five half-lives prior to the use of Adenoscan.

146 Carcinogenesis, Mutagenesis, Impairment of Fertility

- 147 Studies in animals have not been performed to evaluate the carcinogenic potential of Adenoscan
- 148 (adenosine injection). Adenosine was negative for genotoxic potential in the Salmonella (Ames
- 149 Test) and Mammalian Microsome Assay.
- 150 Adenosine, however, like other nucleosides at millimolar concentrations present for several
- 151 doubling times of cells in culture, is known to produce a variety of chromosomal alterations.
- 152 Fertility studies in animals have not been conducted with adenosine.

153 Pregnancy Category C

- 154 Animal reproduction studies have not been conducted with adenosine; nor have studies been
- 155 performed in pregnant women. Because it is not known whether Adenoscan can cause fetal harm
- 156 when administered to pregnant women, Adenoscan should be used during pregnancy only if
- 157 clearly needed.

158 **Pediatric Use**

159 The safety and effectiveness of Adenoscan in patients less than 18 years of age have not been160 established.

161 Geriatric Use

162 Clinical studies of Adenoscan did not include sufficient numbers of subjects aged younger than

163 65 years to determine whether they respond differently. Other reported experience has not

164 revealed clinically relevant differences of the response of elderly in comparison to younger

165 patients. Greater sensitivity of some older individuals, however, cannot be ruled out.

166 **ADVERSE REACTIONS**

167 The following reactions with an incidence of at least 1% were reported with intravenous

168 Adenoscan among 1421 patients enrolled in controlled and uncontrolled U.S. clinical trials.

169 Despite the short half-life of adenosine, 10.6% of the side effects occurred not with the infusion

170 of Adenoscan but several hours after the infusion terminated. Also, 8.4% of the side effects that

171 began coincident with the infusion persisted for up to 24 hours after the infusion was complete.

172 In many cases, it is not possible to know whether these late adverse events are the result of

173 Adenoscan infusion.

Flushing	44%
Chest discomfort	40%
Dyspnea or urge to	28%
breathe deeply	
Headache	18%
Throat, neck or jaw	15%
discomfort	
Gastrointestinal	13%
discomfort	
Lightheadedness/dizziness	12%
Upper extremity	4%
discomfort	
ST segment depression	3%
First-degree AV block	3%
Second-degree AV block	3%
Paresthesia	2%
Hypotension	2%

Nervousness	2%
Arrhythmias	1%

- 174 Adverse experiences of any severity reported in less than 1% of patients include:
- 175 **Body as a whole:** back discomfort; lower extremity discomfort; weakness.
- 176 Cardiovascular System: nonfatal myocardial infarction; life-threatening ventricular arrhythmia;
- 177 third-degree AV block; bradycardia; palpitation; sinus exit block; sinus pause; sweating; T-wave
- 178 changes; hypertension (systolic blood pressure > 200 mm Hg).
- 179 Central Nervous System: drowsiness; emotional instability; tremors.
- 180 Genital/Urinary System: vaginal pressure; urgency.
- 181 **Respiratory System:** cough.
- 182 **Special Senses:** blurred vision; dry mouth; ear discomfort; metallic taste; nasal congestion;
- 183 scotomas; tongue discomfort.

184 **Post Marketing Experience** (see WARNINGS)

- 185 The following adverse events have been reported from marketing experience with Adenoscan.
- 186 Because these events are reported voluntarily from a population of uncertain size, are associated
- 187 with concomitant diseases and multiple drug therapies and surgical procedures, it is not always
- 188 possible to reliably estimate their frequency or establish a causal relationship to drug exposure.
- 189 Decisions to include these events in labeling are typically based on one or more of the following
- 190 factors: (1) seriousness of the event, (2) frequency of the reporting, (3) strength of causal
- 191 connection to the drug, or a combination of these factors.

- **Body as a Whole**
- 194 Injection site reaction
- 195Central Nervous System
- 196 Seizure activity, including tonic clonic (grand mal) seizures, and loss of consciousness
- 197 **Digestive**
- 198 Nausea and vomiting
- 199 **Respiratory**
- 200 Respiratory arrest
- 201

202 **OVERDOSAGE**

- 203 The half-life of adenosine is less than 10 seconds and side effects of Adenoscan (when they
- 204 occur) usually resolve quickly when the infusion is discontinued, although delayed or persistent
- 205 effects have been observed. Methylxanthines, such as caffeine and theophylline, are competitive
- adenosine receptor antagonists and theophylline has been used to effectively terminate persistent
- side effects. In controlled U.S. clinical trials, theophylline (50-125 mg slow intravenous
- 208 injection) was needed to abort Adenoscan side effects in less than 2% of patients.

210 DOSAGE AND ADMINISTRATION

211 For intravenous infusion only.

Adenoscan should be given as a continuous peripheral intravenous infusion.

The recommended intravenous dose for adults is 140 mcg/kg/min infused for six minutes (totaldose of 0.84 mg/kg).

215 The required dose of thallium-201 should be injected at the midpoint of the Adenoscan infusion

216 (i.e., after the first three minutes of Adenoscan). Thallium-201 is physically compatible with

217 Adenoscan and may be injected directly into the Adenoscan infusion set.

218 The injection should be as close to the venous access as possible to prevent and inadvertent

219 increase in the dose of Adenoscan (the contents of the IV tubing) being administered.

220 There are no data on the safety or efficacy of alternative Adenoscan infusion protocols.

The safety and efficacy of Adenoscan administered by the intracoronary route have not beenestablished.

- 223 The following Adenoscan infusion nomogram may be used to determine that appropriate
- 224 infusion rate corrected for total body weight:

Patient Weight		Infusion Rate
kg	lbs	mL/min
45	99	2.1
50	110	2.3
55	121	2.6
60	132	2.8
65	143	3.0

70	154	3.3
75	165	3.5
80	176	3.8
85	187	4.0
90	198	4.2

225 This nomogram was derived from the following general formula:

0.140 (mg/kg/min) x total body weight (kg) Adenoscan concentration (3 mg/mL)

Infusion rate (mL/min)

=

227 **Note:** Parenteral drug products should be inspected visually for particulate matter and

228 discoloration prior to administration.

HOW SUPPLIED

- Adenoscan (adenosine injection) is supplied as 20 mL and 30 mL vials of sterile nonpyrogenic
- 231 solution in normal saline.

Product	NDC No.	
Code		
87120	0469-0871-20	60 mg/20 mL (3 mg/mL) in a 20 mL single-dose, flip-top glass vial, packaged individually and in packages of ten.
87130	0469-0871-30	90 mg/30 mL (3 mg/mL) in a 30 mL single-dose, flip-top glass vial, packaged individually and in packages of

ten.

- 232 Store at controlled room temperature 15°-30°C (59°-86°F)
- 233 Do not refrigerate as crystallization may occur. If crystallization has occurred, dissolve crystals
- by warming at room temperature. The solution must be clear at the time of use.
- 235 Contains no preservative. Discard unused portion.
- 236 **Rx only**
- 237
- 238 Marketed by:
- Astellas Pharma US, Inc.
- 240 Deerfield, IL 60015
- 241 Manufactured by:
- Hospira, Inc.
- 243 Lake Forest, IL 60045 USA
- 244 Revised: April 2005