

SAFETY INSPECTION REPORT AND COMPLIANCE INSPECTION

1. LICENSEE/LOCATION INSPECTED: Kirkwood Medical Group, LLC 3844 South Lindbergh Blvd. Sunset Hills, MO 63127 REPORT NUMBER(S) 2008-001	2. NRC/REGIONAL OFFICE U.S. Nuclear Regulatory Commission Region III 2443 Warrenville Road Suite 210 Lisle, Illinois 60532-4351
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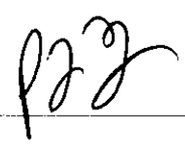
3. DOCKET NUMBER(S) 030-37527	4. LICENSEE NUMBER(S) 24-32667-01	5. DATE(S) OF INSPECTION 5/23/08
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LICENSEE:
The inspection was an examination of the activities conducted under your license as they relate to radiation safety and to compliance with the Nuclear Regulatory Commission (NRC) rules and regulations and the conditions of your license. The inspection consisted of selective examinations of procedures and representative records, interviews with personnel, and observations by the inspector. The inspection findings are as follows:

- 1. Based on the inspection findings, no violations were identified.
- 2. Previous violation(s) closed.
- 3. The violation(s), specifically described to you by the inspector as non-cited violations, are not being cited because they were self-identified, non-repetitive, and corrective action was or is being taken, and the remaining criteria in the NRC Enforcement Policy, NUREG-1600, to exercise discretion, were satisfied.

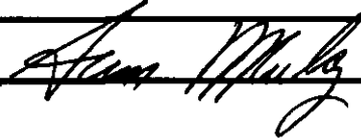
_____ Non-Cited Violation(s) was/were discussed involving the following requirement(s) and Corrective Action(s):

- 4. During this inspection certain of your activities, as described below and/or attached, were in violation of NRC requirements and are being cited. This form is a NOTICE OF VIOLATION, which may be subject to posting in accordance with 10 CFR 19.11.
(Violations and Corrective Actions)



Licensee's Statement of Corrective Actions for Item 4, above.

I hereby state that, within 30 days, the actions described by me to the inspector will be taken to correct the violations identified. This statement of corrective actions is made in accordance with the requirements of 10 CFR 2.201 (corrective steps already taken, corrective steps which will be taken, date when full compliance will be achieved). I understand that no further written response to NRC will be required, unless specifically requested.

Title	Printed Name	Signature	Date
LICENSEE'S REPRESENTATIVE			
NRC INSPECTOR	Sam Mulay		5/23/08

**SAFETY INSPECTION REPORT
AND COMPLIANCE INSPECTION**

1. LICENSEE

Kirkwood Medical Group, LLC
REPORT NUMBER(S)
2008-001

2. NRC/REGIONAL OFFICE

Region III
2443 Warrenville Road, Suite 210
Lisle, IL 60532

3. DOCKET NUMBER(S)

030-37527

4. LICENSE NUMBER(S)

24-32667-01

5. DATE(S) OF INSPECTION

May 23, 2008

6. INSPECTION PROCEDURES USED

87130

7. INSPECTION FOCUS AREAS

03.01-03.07**SUPPLEMENTAL INSPECTION INFORMATION**

1. PROGRAM CODE(S)

2201

2. PRIORITY

G5

3. LICENSEE CONTACT

Sandy Fischer, Office Mgr.

4. TELEPHONE NUMBER

314-447-1301

Main Office Inspection

Next Inspection Date: May 2013

Field Office _____

Temporary Job Site
Inspection _____**PROGRAM SCOPE**

This review was an initial inspection of this cardiology program which became active in November 2007. The licensee performs approximately 12 diagnostic cardiac procedures per week using both technetium-99m card iolite and thallium-201. Currently, one part-time technologist performs all patient procedures on Monday and Tuesday only from 7:30 am to 4:30 pm. Generators are not received and all material is obtained from an area pharmacy in the form of unit doses. An outside consultant performs quarterly program audits which appear to adequately oversee licensed activities.

Performance Observations

Interviews were conducted with the technologist via telephone and revealed an adequate level of understanding of emergency and material handling procedures and techniques. Dose calibrator constancy checks, daily area surveys, package receipt procedures, and patient injection techniques were described with no problems noted.

Licensed material was observed adequately secured upon arrival at the facility and was not readily accessible to members of the general public. Independent measurements taken did not indicate readings in excess of background levels (0.02 mr/hr).

Dosimetry records reviewed indicated whole-body and extremity readings for YTD 2008 of 98 mRem and 340 mRem respectively.