CLIA ANNUAL LABORATORY REGISTRY
2010

Once a year the Centers for Medicare and Medicaid Services makes available to physicians and to the general public specific information (including information provided to CMS by the Office of the Inspector General) that is useful in evaluating the performance of laboratories. The Clinical Laboratory Improvement Amendments of 1988 (CLIA) and implementing regulations at 42 CFR 493.1850 require that this listing include the following:

(1) A list of laboratories that have been convicted, under Federal or State laws relating to fraud and abuse, false billing, or kickbacks.

(2) A list of laboratories that have had their CLIA certificates suspended, limited, or revoked, and the reasons for the adverse actions.

(3) A list of persons who have been convicted of violating CLIA requirements, as specified in section 353(1) of the PHS Act, together with circumstances of each case and the penalties imposed.

(4) A list of laboratories on which alternative sanctions have been imposed, showing--
   (i) the effective date of the sanctions;
   (ii) the reason for imposing them;
   (iii) any corrective action taken by the laboratory;
   (iv) if the laboratory has achieved compliance, the verified date of compliance.

(5) A list of laboratories whose accreditation has been withdrawn or revoked and the reasons for the withdrawal or revocation.

(6) All appeals and hearing decisions.

(7) A list of laboratories against which CMS has brought suit under Section 493.1846 and the reasons for those actions.

(8) A list of laboratories that have been excluded from participation in Medicare or Medicaid and the reasons for exclusion.

Civil settlements reached with clinical laboratories are also noted.

The Laboratory Registry is compiled for the calendar year proceeding the date the information is made available and also contains corrections of any erroneous statements of information that appeared in the previous registry. A final section includes other specific information that may be useful in evaluating the performance of laboratories, as specified in 42 CFR 493.1850(a). It also includes information provided by CLIA exempt states.
JOWHER KHALEEL, DIRECTOR
JOWHER KHALEEL MD PC
20000 FARMINGTON ROAD BUILDING E
LIVONIA, MI 48154
CLIA ID# 23D0941245

SANCTION: CANCEL MEDICARE/MEDICAID APPROVAL
SUSPENSION OF CLIA CERTIFICATE
REVOCATION OF CLIA CERTIFICATE

EFFECTIVE DATE: AUGUST 18, 2010

REASON: CONDITION LEVEL NONCOMPLIANCE

REGINALD SHARPE DO, DIRECTOR
WOMEN'S ADVISORY CENTER
43700 N WOODWARD, SUITE 104
BLOOMFIELD HILLS, MI 48302
CLIA ID# 23D0979850

SANCTION: CANCEL MEDICARE/MEDICAID APPROVAL
LIMITATION OF CLIA CERTIFICATE
REVOCATION OF CLIA CERTIFICATE

EFFECTIVE DATE: DECEMBER 17, 2010

REASON: CONDITION LEVEL NONCOMPLIANCE

GREGORY MC NAMARA, DIRECTOR
FAMILY MEDICAL CENTER
811 SE SECOND STREET SUITE A
LITTLE FALLS, MN 56345
CLIA ID# 24D0405950

SANCTION: CANCEL MEDICARE/MEDICAID APPROVAL
REVOCATION OF CLIA CERTIFICATE

EFFECTIVE DATE: OCTOBER 07, 2010

REASON: IMPROPER PROFICIENCY TESTING REFERRAL.
SANCTION: DIRECTED PLAN OF CORRECTION
SUSPENSION PART OF MEDICARE/MEDICAID

EFFECTIVE DATE: DECEMBER 17, 2010

REASON: CANCELLATION OF MEDICARE APPROVAL REVOCATION

KEVIN CAVANAGH, DIRECTOR
HEALTH DEPT OF NORTHWEST MI - NORTHERN MICHIGAN
REGIONAL LABORATORY
95 LIVINGSTON BLVD SUITE D
GAYLORD, MI 49735
CLIA ID# 23D1038040

SANCTION: DIRECTED PLAN OF CORRECTION

EFFECTIVE DATE: SEPTEMBER 21, 2010

REASON: CONDITION LEVEL NONCOMPLIANCE

STATUS: COMPLIANCE ACHIEVED

MARK ROBIA, DIRECTOR
NORTH COUNTRY REGIONAL HOSPITAL LAB
1300 ANNE STREET NW
BEMIDJI, MN 56601
CLIA ID# 24D0041382

SANCTION: DIRECTED PLAN OF CORRECTION

EFFECTIVE DATE: FEBRUARY 22, 2010

REASON: CONDITION LEVEL NONCOMPLIANCE

STATUS: COMPLIANCE ACHIEVED