### §493.933

value has been established for each response, the appropriateness of the response must be determined by using either fixed criteria based on the percentage difference from the target value or the number of standard deviations (SDs) the response differs from the target value.

Criteria for Acceptable Performance

The criteria for acceptable performance are—

| Analyte or test                           | Criteria for acceptable per-<br>formance      |
|---|---|
| Alanine aminotransferase<br>(ALT/SGPT).   | Target value ±20%.                            |
| Albumin                                   | Target value ±10%.                            |
| Alkaline phosphatase                      | Target value ±30%.                            |
| Amylase                                   | Target value ±30%.                            |
| Aspartate aminotransferase<br>(AST/SGOT). | Target value ±20%.                            |
| Bilirubin, total                          | Target value ±0.4 mg/dL or<br>±20% (greater). |
| Blood gas pO2                             | Target value ±3 SD.                           |
| pCO2                                      | Target value ±5 mm Hg or<br>±8% (greater).    |
| рН  | Target value ±0.04.                           |
| Calcium, total                            | Target value ±1.0 mg/dL.                      |
| Chloride                                  | Target value ±5%.                             |
| Cholesterol, total                        | Target value ±10%.                            |
| Cholesterol, high density lipoprotein.    | Target value ±30%.                            |

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| Analyte or test  | Criteria for acceptable per-<br>formance                        |
|--|---|
| Creatine kinase  | Target value ±30%.  |
| Creatine kinase isoenzymes   | MB elevated (presence or ab-<br>sence) or Target value<br>±3SD. |
| Creatinine   | Target value ±0.3 mg/dL or ±15% (greater).                      |
| Glucose (excluding glucose<br>performed on monitoring<br>devices cleared by FDA for<br>home use. | Target value ±6 mg/dl or<br>±10% (greater).                     |
| Iron, total  | Target value ±20%.  |
| Lactate dehydrogenase (LDH).   | Target value ±20%.  |
| LDH isoenzymes   | LDH1/LDH2 (+ or -) or Tar-<br>get value ±30%.                   |
| Magnesium  | Target value ±25%.  |
| Potassium  | Target value ±0.5 mmol/L.                                       |
| Sodium   | Target value ±4 mmol/L.   |
| Total Protein  | Target value ±10%.  |
| Triglycerides  | Target value ±25%.  |
| Urea nitrogen  | Target value ±2 mg/dL or<br>±9% (greater).                      |
| Uric acid  | Target value ±17%.  |

(3) The criterion for acceptable performance for qualitative routine chemistry tests is positive or negative.

(4) To determine the analyte testing event score, the number of acceptable analyte responses must be averaged using the following formula:

| Number of acceptable responses for the analyte $\times 100 =$ | Analyte score for |
|---|-------------------|
| Total number of challenges for the analyte                    | the testing event |

(5) To determine the overall testing event score, the number of correct responses for all analytes must be averaged using the following formula:

 $\frac{\text{Number of acceptable responses for all challenges}}{\text{Total number of all challenges}} \times 100 = \text{Testing event score}$ 

[57 FR 7151, Feb. 28, 1992, as amended at 68 FR 3702, Jan. 24, 2003]

## §493.933 Endocrinology.

(a) Program content and frequency of challenge. To be approved for proficiency testing for endocrinology, a program must provide a minimum of five samples per testing event. There must be at least three testing events at approximately equal intervals per year. The annual program must provide samples that cover the clinically relevant range of values that would be expected in patient specimens. The samples may be provided through mailed shipments or, at HHS' option, may be provided to HHS or its designee for on-site testing.

(b) *Challenges per testing event*. The minimum number of challenges per testing event a program must provide for each analyte or test procedure is five serum, plasma, blood, or urine samples.

#### Centers for Medicare & Medicaid Services, HHS

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Analyte or Test

Cortisol

Free Thyroxine

Human Chorionic gonadotropin (excluding urine pregnancy tests done by visual color comparison categorized as waived tests) T3 Ubtake

Triiodothyronine

Thyroid-stimulating hormone

Thyroxine

(c) Evaluation of a laboratory's analyte or test performance. HHS approves only those programs that assess the accuracy of a laboratory's responses in accordance with paragraphs (c)(1) through (5) of this section.

(1) To determine the accuracy of a laboratory's response for qualitative and quantitative endocrinology tests or analytes, a program must compare the laboratory's response for each analyte with the response that reflects agreement of either 80 percent of ten or more referee laboratories or 80 percent or more of all participating laboratories. The score for a sample in endocrinology is either the score determined under paragraph (c)(2) or (c)(3) of this section.

(2) For quantitative endocrinology tests or analytes, the program must determine the correct response for each analyte by the distance of the response from the target value. After the target value has been established for each response, the appropriateness of the response must be determined by using either fixed criteria based on the percentage difference from the target value or the number of standard deviations (SDs) the response differs from the target value.

Criteria for Acceptable Performance

The criteria for acceptable performance are—

| Analyte or test  | Criteria for acceptable per-<br>formance   |
|--|--|
| Cortisol<br>Free Thyroxine<br>Human Chorionic<br>Gonadotropin (excluding<br>urine pregnancy tests done<br>by visual color comparison<br>categorized as waived<br>tests). | Target value ±25%.<br>Target value ±3 SD.<br>Target value ±3 SD positive<br>or negative.                           |
| T3 Uptake<br>Triiodothyronine<br>Thyroid-stimulating hormone<br>Thyroxine  | Target value ±3 SD.<br>Target value ±3 SD.<br>Target value ±3 SD.<br>Target value ±20% or 1.0<br>mcg/dL (greater). |

(3) The criterion for acceptable performance for qualitative endocrinology tests is positive or negative.

(4) To determine the analyte testing event score, the number of acceptable analyte responses must be averaged using the following formula:

# $\frac{\text{Number of acceptable responses for the analyte}}{\text{Total number of challenges for the analyte}} \times 100 = \frac{\text{Analyte score for the testing event}}{\text{the testing event}}$

(5) To determine the overall testing event score, the number of correct responses for all analytes must be averaged using the following formula:

Number of acceptable responses for all challenges  $\times 100 =$  Testing event score

## Total number of all challenges

[57 FR 7151, Feb. 28, 1992, as amended at 58 FR 5229, Jan. 19, 1993; 68 FR 3702, Jan. 24, 2003]

#### §493.937 Toxicology.

(a) *Program content and frequency of challenge.* To be approved for proficiency testing for toxicology, the annual program must provide a minimum of five samples per testing event. There