

Food and Drug Administration, HHS

§ 862.3

862.3040	Alcohol test system.
862.3050	Breath-alcohol test system.
862.3100	Amphetamine test system.
862.3110	Antimony test system.
862.3120	Arsenic test system.
862.3150	Barbiturate test system.
862.3170	Benzodiazepine test system.
862.3200	Clinical toxicology calibrator.
862.3220	Carbon monoxide test system.
862.3240	Cholinesterase test system.
862.3250	Cocaine and cocaine metabolite test system.
862.3270	Codeine test system.
862.3280	Clinical toxicology control material.
862.3300	Digitoxin test system.
862.3320	Digoxin test system.
862.3350	Diphenylhydantoin test system.
862.3380	Ethosuximide test system.
862.3450	Gentamicin test system.
862.3520	Kanamycin test system.
862.3550	Lead test system.
862.3555	Lidocaine test system.
862.3560	Lithium test system.
862.3580	Lysergic acid diethylamide (LSD) test system.
862.3600	Mercury test system.
862.3610	Methamphetamine test system.
862.3620	Methadone test system.
862.3630	Methaqualone test system.
862.3640	Morphine test system.
862.3645	Neuroleptic drugs radioreceptor assay test system.
862.3650	Opiate test system.
862.3660	Phenobarbital test system.
862.3670	Phenothiazine test system.
862.3680	Primidone test system.
862.3700	Propoxyphene test system.
862.3750	Quinine test system.
862.3830	Salicylate test system.
862.3850	Sulfonamide test system.
862.3870	Cannabinoid test system.
862.3880	Theophylline test system.
862.3900	Tobramycin test system.
862.3910	Tricyclic antidepressant drugs test system.
862.3950	Vancomycin test system.

AUTHORITY: 21 U.S.C. 351, 360, 360c, 360e, 360j, 371.

SOURCE: 52 FR 16122, May 1, 1987, unless otherwise noted.

Subpart A—General Provisions

§ 862.1 Scope.

(a) This part sets forth the classification of clinical chemistry and clinical toxicology devices intended for human use that are in commercial distribution.

(b) The identification of a device in a regulation in this part is not a precise description of every device that is, or will be, subject to the regulation. A

manufacturer who submits a pre-market notification submission for a device under part 807 cannot show merely that the device is accurately described by the section title and identification provisions of a regulation in this part, but shall state why the device is substantially equivalent to other devices, as required in § 807.87.

(c) References in this part to regulatory sections of the Code of Federal Regulations are to chapter I of title 21 unless otherwise noted.

(d) Guidance documents referenced in this part are available on the Internet at <http://www.fda.gov/cdrh/guidance.html>.

[52 FR 16122, May 1, 1987, as amended at 67 FR 58329, Sept. 16, 2002]

§ 862.2 Regulation of calibrators.

Many devices classified in this part are intended to be used with a calibrator. A calibrator has a reference value assigned to it which serves as the basis by which test results of patients are derived or calculated. The calibrator for a device may be (a) manufactured and distributed separately from the device with which it is intended to be used, (b) manufactured and distributed as one of several device components, such as in a kit of reagents, or (c) built-in as an integral part of the device. Because of the central role that a calibrator plays in the measurement process and the critical effect calibrators have on accuracy of test results, elsewhere in this part, all three of these types of calibrators (§§ 862.1150 and 862.3200 of this part) are classified into class II, notwithstanding the classification of the device with which it is intended to be used. Thus, a device and its calibrator may have different classifications, even if the calibrator is built into the device.

§ 862.3 Effective dates of requirement for premarket approval.

A device included in this part that is classified into class III (premarket approval) shall not be commercially distributed after the date shown in the regulation classifying the device unless the manufacturer has an approval under section 515 of the act (unless an exemption has been granted under section 520(g)(2) of the act). An approval