

AMENDED IN SENATE MAY 21, 2009

AMENDED IN SENATE MAY 14, 2009

AMENDED IN SENATE APRIL 22, 2009

SENATE BILL

No. 744

Introduced by Senator Strickland

February 27, 2009

An act to amend Sections 1206, 1223, 1246, 1300, 1301, and 1302 of, and to add Section 1300.2 to, the Business and Professions Code, relating to clinical laboratories, ~~making an appropriation therefor~~, and declaring the urgency thereof, to take effect immediately.

LEGISLATIVE COUNSEL'S DIGEST

SB 744, as amended, Strickland. Clinical laboratories.

(1) Existing law provides for the licensure, registration, and regulation of clinical laboratories and various clinical laboratory personnel by the State Department of Public Health and makes a violation of those provisions a crime.

Existing law requires the department to deem certain laboratories accredited by private, nonprofit organizations as meeting state licensure or registration requirements if certain conditions are met. Under existing law, the private, nonprofit organization must, among other things, be approved by the Health Care Financing Administration (HCFA) of the federal Department of Health and Human Services and must be approved by the department as having accreditation standards that are equal to, or more stringent than, state requirements for licensure or registration. The laboratory must meet the accreditation standards of that organization and must agree to permit the organization to provide records or other information to the department.

This bill would require the private, nonprofit organization to be approved by the federal Center for Medicare and Medicaid Services instead of HCFA, to conduct inspections of clinical laboratories in a manner that will determine compliance with existing law, as specified, and to provide the department with additional information including, among other things, a detailed description of the inspection process and a description of the process for monitoring proficiency testing performance. The bill would also require the laboratory to meet additional conditions, including authorizing the private, nonprofit organization to release specified performance testing results and notification of condition-level requirement violations or withdrawal of laboratory accreditation.

Existing law specifies various fees applicable to clinical laboratories and laboratory personnel and requires the deposit of those fees in the Clinical Laboratory Improvement Fund. Existing law requires that, upon appropriation, moneys deposited in that fund be expended by the department to administer these provisions. Existing law requires the issuance of a separate license for each laboratory location, except as specified. Among other entities, not-for-profit, or federal, state, or local government laboratories engaging in limited public health testing are authorized to apply for a single license or registration, as specified.

This bill would impose a fee for approval of each of those laboratories and would increase certain other fees applicable to laboratories and laboratory personnel. The bill would prohibit the fees imposed from exceeding the costs incurred by the department in regulating clinical laboratories and their personnel. The bill ~~also would provide that moneys deposited in the Clinical Laboratory Improvement Fund are continuously appropriated to the department for specified purposes, thereby making an appropriation, and~~ *would require all interest earned on moneys deposited in the Clinical Laboratory Improvement Fund to be maintained in the fund and* would prohibit the redirection of ~~those moneys in the fund~~ for any other purpose. The bill would require the department, ~~beginning in fiscal year 2013–14, to report to the Legislature by July 1, 2013, on the status of its work toward receiving an exemption from federal CLIA oversight~~ *extent to which the state clinical laboratory oversight program meets or exceeds federal standards, the extent to which the federal government is accepting exemption applications from states relative to federal CLIA oversight, and the potential cost to the state for an exemption.*

Existing law provides for the renewal of a clinical laboratory license or registration and requires that the renewal fee be paid during the 30-day period before the expiration of the license or registration. Existing law specifies that failure to pay the renewal fee results in forfeiture of the license or registration after a period of 60 days from the expiration date.

This bill would require a licensee or registrant that fails to renew a license or registration before the expiration date to pay a specified delinquency fee for up to 60 days after the expiration date, in addition to the renewal fee.

(2) This bill would declare that it is to take effect immediately as an urgency statute.

Vote: $\frac{2}{3}$. Appropriation: ~~yes~~-no. Fiscal committee: yes.
State-mandated local program: no.

The people of the State of California do enact as follows:

1 ~~SECTION 1. Consistent with the intent of Senate Bill 113 of~~
2 ~~the 1995–96 Regular Session (Chapter 510, Statutes of 1995), it~~
3 ~~is the intent of the Legislature, by this act, to enact provisions in~~
4 ~~state law that ensure that the state has enforcement authority and~~
5 ~~administrative structures adequate to enforce the requirements of~~
6 ~~the federal Clinical Laboratory Improvement Amendments of 1988~~
7 ~~(CLIA) and state laws regulating clinical laboratories, and that~~
8 ~~will enable the state to seek approval from the United States~~
9 ~~Secretary of Health and Human Services to exempt from CLIA~~
10 ~~those clinical laboratories licensed or registered under state law.~~

11 ~~SEC. 2.~~

12 SECTION 1. Section 1206 of the Business and Professions
13 Code is amended to read:

14 1206. (a) For the purposes of this chapter the following
15 definitions are applicable:

16 (1) “Biological specimen” means any material that is derived
17 from the human body.

18 (2) “Blood electrolyte analysis” means the measurement of
19 electrolytes in a blood specimen by means of ion selective
20 electrodes on instruments specifically designed and manufactured
21 for blood gas and acid-base analysis.

(3) “Blood gas analysis” means a clinical laboratory test or examination that deals with the uptake, transport, and metabolism of oxygen and carbon dioxide in the human body.

(4) “Clinical laboratory test or examination” means the detection, identification, measurement, evaluation, correlation, monitoring, and reporting of any particular analyte, entity, or substance within a biological specimen for the purpose of obtaining scientific data which may be used as an aid to ascertain the presence, progress, and source of a disease or physiological condition in a human being, or used as an aid in the prevention, prognosis, monitoring, or treatment of a physiological or pathological condition in a human being, or for the performance of nondiagnostic tests for assessing the health of an individual.

(5) “Clinical laboratory science” means any of the sciences or scientific disciplines used to perform a clinical laboratory test or examination.

(6) “Clinical laboratory practice” means the application of clinical laboratory sciences or the use of any means that applies the clinical laboratory sciences within or outside of a licensed or registered clinical laboratory. Clinical laboratory practice includes consultation, advisory, and other activities inherent to the profession.

(7) “Clinical laboratory” means any place used, or any establishment or institution organized or operated, for the performance of clinical laboratory tests or examinations or the practical application of the clinical laboratory sciences. That application may include any means that applies the clinical laboratory sciences.

(8) “Direct and constant supervision” means personal observation and critical evaluation of the activity of unlicensed laboratory personnel by a physician and surgeon, or by a person licensed under this chapter other than a trainee, during the entire time that the unlicensed laboratory personnel are engaged in the duties specified in Section 1269.

(9) “Location” means either a street and city address, or a site or place within a street and city address, where any of the clinical laboratory sciences or scientific disciplines are practiced or applied, or where any clinical laboratory tests or examinations are performed.

(10) “Physician office laboratory” means a clinical laboratory that is licensed or registered under Section 1265, and that is either: (A) a clinical laboratory that is owned and operated by a partnership or professional corporation that performs clinical laboratory tests or examinations only for patients of five or fewer physicians and surgeons or podiatrists who are shareholders, partners, or employees of the partnership or professional corporation that owns and operates the clinical laboratory; or (B) a clinical laboratory that is owned and operated by an individual licensed physician and surgeon or a podiatrist, and that performs clinical laboratory tests or examinations only for patients of the physician and surgeon or podiatrist who owns and operates the clinical laboratory.

(11) “Public health laboratory” means a laboratory that is operated by a city or county in conformity with Article 5 (commencing with Section 101150) of Chapter 2 of Part 3 of Division 101 of the Health and Safety Code and the regulations adopted thereunder.

(12) “Specialty” means histocompatibility, microbiology, diagnostic immunology, chemistry, hematology, immunochemistry, pathology, genetics, or other specialty specified by regulation adopted by the department.

(13) “Subspecialty” for purposes of microbiology, means bacteriology, mycobacteriology, mycology, parasitology, virology, molecular biology, and serology for diagnosis of infectious diseases, or other subspecialty specified by regulation adopted by the department; for purposes of diagnostic immunology, means syphilis serology, general immunology, or other subspecialty specified by regulation adopted by the department; for purposes of chemistry, means routine chemistry, clinical microscopy, endocrinology, toxicology, or other subspecialty specified by regulation adopted by the department; for purposes of immunochemistry, means ABO/Rh Type and Group, antibody detection for transfusion, antibody detection nontransfusion, antibody identification, compatibility, or other subspecialty specified by regulation adopted by the department; for pathology, means tissue pathology, oral pathology, diagnostic cytology, or other subspecialty specified by regulation adopted by the department; for purposes of genetics, means molecular biology related to the diagnosis of human genetic abnormalities,

1 cytogenetics, or other subspecialty specified by regulation adopted
2 by the department.

3 (14) “Direct and responsible supervision” means both of the
4 following:

5 (A) Personal observation and critical evaluation of the activity
6 of a trainee by a physician and surgeon, or by a person licensed
7 under this chapter other than a trainee, during the entire time that
8 the trainee is performing clinical laboratory tests or examinations.

9 (B) Personal review by the physician and surgeon or the licensed
10 person of all results of clinical laboratory testing or examination
11 performed by the trainee for accuracy, reliability, and validity
12 before the results are reported from the laboratory.

13 (15) “Licensed laboratory” means a clinical laboratory licensed
14 pursuant to paragraph (1) of subdivision (a) of Section 1265.

15 (16) “Registered laboratory” means a clinical laboratory
16 registered pursuant to paragraph (2) of subdivision (a) of Section
17 1265.

18 (17) “Point-of-care laboratory testing device” means a portable
19 laboratory testing instrument to which the following applies:

20 (A) It is used within the proximity of the patient for whom the
21 test or examination is being conducted.

22 (B) It is used in accordance with the patient test management
23 system, the quality control program, and the comprehensive quality
24 assurance program established and maintained by the laboratory
25 pursuant to paragraph (2) of subdivision (d) of Section 1220.

26 (C) It meets the following criteria:

27 (i) Performs clinical laboratory tests or examinations classified
28 as waived or of moderate complexity under CLIA.

29 (ii) Performs clinical laboratory tests or examinations on
30 biological specimens that require no preparation after collection.

31 (iii) Provides clinical laboratory tests or examination results
32 without calculation or discretionary intervention by the testing
33 personnel.

34 (iv) Performs clinical laboratory tests or examinations without
35 the necessity for testing personnel to perform calibration or
36 maintenance, except resetting pursuant to the manufacturer’s
37 instructions or basic cleaning.

38 (18) “Analyte” means the substance or constituent being
39 measured including, but not limited to, glucose, sodium, or
40 theophylline, or any substance or property whose presence or

1 absence, concentration, activity, intensity, or other characteristics
2 are to be determined.

3 (b) Nothing in this chapter shall restrict, limit, or prevent any
4 person licensed to provide health care services under the laws of
5 this state, including, but not limited to, licensed physicians and
6 surgeons and registered nurses, from practicing the profession or
7 occupation for which he or she is licensed.

8 (c) Nothing in this chapter shall authorize any person to perform
9 or order health care services, or utilize the results of the clinical
10 laboratory test or examination, unless the person is otherwise
11 authorized to provide that care or utilize the results. The inclusion
12 of a person in Section 1206.5 for purposes of performing a clinical
13 laboratory test or examination shall not be interpreted to authorize
14 a person, who is not otherwise authorized, to perform venipuncture,
15 arterial puncture, or skin puncture.

16 ~~SEC. 3.~~

17 *SEC. 2.* Section 1223 of the Business and Professions Code is
18 amended to read:

19 1223. (a) The Legislature finds and declares that it is the public
20 policy of the state to ensure that California's laboratory standards,
21 including its laboratory personnel standards, be sustained in order
22 to provide accurate, reliable, and necessary test results. The
23 Legislature further finds that inspections are the most effective
24 means of furthering this policy. It is not the intent of the Legislature
25 to reduce in any way the resources available to the department for
26 inspections, but rather to provide the department with the greatest
27 flexibility to concentrate its resources where they can be most
28 effective. It is the intent of the Legislature to provide for an
29 inspection process that includes state-based inspection components
30 and that determines compliance with federal and state requirements
31 for clinical laboratories.

32 (b) The department shall employ, or contract for, inspectors,
33 special agents, and investigators, and provide any clerical and
34 technical assistance as necessary to administer this chapter and
35 may incur other expenses as necessary.

36 (c) Laboratories accredited by a private, nonprofit organization
37 shall be deemed by the department to meet state licensure or
38 registration requirements, and shall be issued a certificate of that
39 deemed status by the department, provided that both of the
40 following conditions are met:

1 (1) The private, nonprofit organization meets all of the following
2 requirements:
3 (A) Is approved by the federal Center for Medicare and Medicaid
4 Services as an accreditation body under CLIA and provides the
5 department with the following information:
6 (i) A detailed comparison of the individual accreditation or
7 approval requirements, with the comparable condition-level
8 requirements.
9 (ii) A detailed description of its inspection process, including
10 all of the following:
11 (I) Frequency of inspections.
12 (II) Copies of inspection forms.
13 (III) Instructions and guidelines.
14 (IV) A description of the review and decisionmaking process
15 of inspections.
16 (V) A statement concerning whether inspections are announced
17 or unannounced.
18 (VI) A description of the steps taken to monitor the correction
19 of deficiencies.
20 (iii) A description of the process for monitoring proficiency
21 testing performance, including action to be taken in response to
22 unsuccessful participation.
23 (iv) A list of all of its current California licensed or registered
24 laboratories and the expiration date of their accreditation, licensure,
25 or registration, as applicable.
26 (v) Procedures for making proficiency testing information
27 available, including explanatory information required to interpret
28 proficiency testing results, on a reasonable basis, upon request of
29 any person.
30 (B) Is approved by the department as having accreditation
31 standards that are equal to, or more stringent than, state
32 requirements for licensure and registration.
33 (C) Conducts inspections of clinical laboratories in a manner
34 that will determine compliance with federal standards and
35 California laws to the extent that California laws provide greater
36 protection to residents, or are more stringent than federal standards,
37 as determined by the department. Notwithstanding any other
38 provision of law, the department may, without taking regulatory
39 action pursuant to Chapter 3.5 (commencing with Section 11340)
40 of Part 1 of Division 3 of Title 2 of the Government Code,

1 implement or interpret this section by means of an All Clinical
2 Laboratories Letter (ACLL). The department shall post the ACLL
3 on its Internet Web site so that any person may observe which
4 California laws are more stringent than federal standards, and
5 which accreditation bodies have been approved to conduct
6 inspections. Public comment on the ACLL shall be accepted by
7 the department for 30 days after posting and shall become final
8 45 days after the posting. Comments received shall be considered
9 by the department. Nothing in this subdivision is intended to
10 change existing statutory or regulatory requirements governing
11 the operation of clinical laboratories or their personnel.

12 (D) Agrees to permit the department or its agents or contractors
13 to conduct random inspections of clinical laboratories accredited
14 by it in order to validate compliance with California law.

15 (2) The laboratory meets all of the following requirements:

16 (A) Meets the accreditation standards of the private, nonprofit
17 organization.

18 (B) Agrees to permit the private, nonprofit organization to
19 provide any records or other information to the department, its
20 agents, or contractors, as the department may require.

21 (C) Pays the applicable fees required under Section 1300.

22 (D) Authorizes its proficiency testing organization to furnish
23 to the private, nonprofit organization the results of the laboratory's
24 participation in an approved proficiency testing program for the
25 purpose of monitoring the laboratory's proficiency testing, along
26 with explanatory information needed to interpret the proficiency
27 testing results, upon request of the department.

28 (E) Authorizes the private, nonprofit organization to release to
29 the department the laboratory's proficiency test results that
30 constitute unsuccessful participation in an approved proficiency
31 testing program, as defined in 42 C.F.R. 493.2, when the laboratory
32 has failed to achieve successful participation in an approved
33 proficiency testing program.

34 (F) Authorizes the private, nonprofit organization to release to
35 the department a notification of every violation of condition-level
36 requirements, including the actions taken by the organization as a
37 result of the violation, within 30 days of the initiation of the action.

38 (G) Authorizes the private, nonprofit organization to give notice
39 to the department of any withdrawal of the laboratory's
40 accreditation.

(d) If the private, nonprofit organization described in subdivision (c) has withdrawn or revoked its accreditation of a laboratory, the laboratory shall retain its certificate of accreditation for 45 days after the laboratory receives notice of the withdrawal or revocation of the accreditation, or the effective date of any action taken by the department, whichever is earlier.

(e) A certificate of deemed status issued pursuant to subdivision (c) shall be renewed annually provided that the conditions for issuance specified in subdivision (c) are still met. Each application for a certificate of deemed status issued under subdivision (c) and each request for renewal of that certificate shall be accompanied by the fees set forth in Section 1300. The total of those certificate application and renewal fees collected by the department shall be sufficient to cover the cost of issuing the certificate. If the department determines that those certificate fees do not fully support the costs of these activities, it shall report that determination to the Legislature.

(f) Nothing in this section shall be construed to prohibit the exercise of the department's authority to conduct complaint investigations, sample validation inspections, or require submission of proficiency testing results to the department to ensure compliance of any clinical laboratory with state standards.

~~SEC. 4.~~

SEC. 3. Section 1246 of the Business and Professions Code is amended to read:

1246. (a) Except as provided in subdivisions (b) and (c), and in Section 23158 of the Vehicle Code, an unlicensed person employed by a licensed clinical laboratory may perform venipuncture or skin puncture for the purpose of withdrawing blood or for clinical laboratory test purposes upon specific authorization from a licensed physician and surgeon provided that he or she meets both of the following requirements:

(1) He or she works under the supervision of a person licensed under this chapter or of a licensed physician and surgeon or of a licensed registered nurse. A person licensed under this chapter, a licensed physician or surgeon, or a registered nurse shall be physically available to be summoned to the scene of the venipuncture within five minutes during the performance of those procedures.

(2) He or she has been trained by a licensed physician and surgeon or by a clinical laboratory bioanalyst in the proper procedure to be employed when withdrawing blood in accordance with training requirements established by the State Department of Public Health and has a statement signed by the instructing physician and surgeon or by the instructing clinical laboratory bioanalyst that the training has been successfully completed.

(b) (1) On and after the effective date of the regulations specified in paragraph (2), any unlicensed person employed by a clinical laboratory performing the duties described in this section shall possess a valid and current certification as a certified phlebotomy technician issued by the department. However, an unlicensed person employed by a clinical laboratory to perform these duties pursuant to subdivision (a) on that date shall have until January 1, 2007, to comply with this requirement, provided that he or she has submitted the application to the department on or before July 1, 2006.

(2) The department shall adopt regulations for certification by January 1, 2001, as a certified phlebotomy technician that shall include all of the following:

(A) The applicant shall hold a valid, current certification as a phlebotomist issued by a national accreditation agency approved by the department, and shall submit proof of that certification when applying for certification pursuant to this section.

(B) The applicant shall complete education, training, and experience requirements as specified by regulations that shall include, but not be limited to, the following:

(i) At least 40 hours of didactic instruction.

(ii) At least 40 hours of practical instruction.

(iii) At least 50 successful venipunctures.

However, an applicant who has been performing these duties pursuant to subdivision (a) may be exempted from the requirements specified in clauses (ii) and (iii), and from 20 hours of the 40 hours of didactic instruction as specified in clause (i), if he or she has at least 1,040 hours of work experience, as specified in regulations adopted by the department.

It is the intent of the Legislature to permit persons performing these duties pursuant to subdivision (a) to use educational leave provided by their employers for purposes of meeting the requirements of this section.

1 (3) Each certified phlebotomy technician shall complete at least
2 three hours per year or six hours every two years of continuing
3 education or training. The department shall consider a variety of
4 programs in determining the programs that meet the continuing
5 education or training requirement.

6 (4) He or she has been found to be competent in phlebotomy
7 by a licensed physician and surgeon or person licensed pursuant
8 to this chapter.

9 (5) He or she works under the supervision of a licensed
10 physician and surgeon, licensed registered nurse, or person licensed
11 under this chapter, or the designee of a licensed physician and
12 surgeon or the designee of a person licensed under this chapter.

13 (6) The department shall adopt regulations establishing standards
14 for approving training programs designed to prepare applicants
15 for certification pursuant to this section. The standards shall ensure
16 that these programs meet the state's minimum education and
17 training requirements for comparable programs.

18 (7) The department shall adopt regulations establishing standards
19 for approving national accreditation agencies to administer
20 certification examinations and tests pursuant to this section.

21 (8) The department shall charge fees for application for and
22 renewal of the certificate authorized by this section of no more
23 than one hundred dollars (\$100) for a two-year period.

24 (c) (1) (A) A certified phlebotomy technician may perform
25 venipuncture or skin puncture to obtain a specimen for
26 nondiagnostic tests assessing the health of an individual, for
27 insurance purposes, provided that the technician works under the
28 general supervision of a physician and surgeon licensed under
29 Chapter 5 (commencing with Section 2000). The physician and
30 surgeon may delegate the general supervision duties to a registered
31 nurse or a person licensed under this chapter, but shall remain
32 responsible for ensuring that all those duties and responsibilities
33 are properly performed. The physician and surgeon shall make
34 available to the department, upon request, records maintained
35 documenting when a certified phlebotomy technician has
36 performed venipuncture or skin puncture pursuant to this
37 paragraph.

38 (B) As used in this paragraph, general supervision requires the
39 supervisor of the technician to determine that the technician is
40 competent to perform venipuncture or skin puncture prior to the

1 technician's first blood withdrawal, and on an annual basis
2 thereafter. The supervisor is also required to determine, on a
3 monthly basis, that the technician complies with appropriate
4 venipuncture or skin puncture policies and procedures approved
5 by the medical director and required by state regulations. The
6 supervisor, or another designated licensed physician and surgeon,
7 registered nurse, or person licensed under this chapter, shall be
8 available for consultation with the technician, either in person or
9 through telephonic or electronic means, at the time of blood
10 withdrawal.

11 (2) (A) Notwithstanding any other provision of law, a person
12 who has been issued a certified phlebotomy technician certificate
13 pursuant to this section may draw blood following policies and
14 procedures approved by a physician and surgeon licensed under
15 Chapter 5 (commencing with Section 2000), appropriate to the
16 location where the blood is being drawn and in accordance with
17 state regulations. The blood collection shall be done at the request
18 and in the presence of a peace officer for forensic purposes in a
19 jail, law enforcement facility, or medical facility, with general
20 supervision.

21 (B) As used in this paragraph, "general supervision" means that
22 the supervisor of the technician is licensed under this code as a
23 physician and surgeon, physician assistant, clinical laboratory
24 bioanalyst, registered nurse, or clinical laboratory scientist, and
25 reviews the competency of the technician before the technician
26 may perform blood withdrawals without direct supervision, and
27 on an annual basis thereafter. The supervisor is also required to
28 review the work of the technician at least once a month to ensure
29 compliance with venipuncture policies, procedures, and regulations.
30 The supervisor, or another person licensed under this code as a
31 physician and surgeon, physician assistant, clinical laboratory
32 bioanalyst, registered nurse, or clinical laboratory scientist, shall
33 be accessible to the location where the technician is working to
34 provide onsite, telephone, or electronic consultation, within 30
35 minutes when needed.

36 (d) The department may adopt regulations providing for the
37 issuance of a certificate to an unlicensed person employed by a
38 clinical laboratory authorizing only the performance of skin
39 punctures for test purposes.

1 ~~SEC. 5.~~

2 *SEC. 4.* Section 1300 of the Business and Professions Code is
3 amended to read:

4 1300. The amount of application, registration, and license fees
5 under this chapter shall be as follows:

6 (a) The application fee for a histocompatibility laboratory
7 director's, clinical laboratory bioanalyst's, clinical chemist's,
8 clinical microbiologist's, clinical laboratory toxicologist's, clinical
9 cytogeneticist's, or clinical molecular biologist's license is
10 sixty-three dollars (\$63) commencing on July 1, 1983.

11 (b) The annual renewal fee for a histocompatibility laboratory
12 director's, clinical laboratory bioanalyst's, clinical chemist's,
13 clinical microbiologist's, or clinical laboratory toxicologist's
14 license is sixty-three dollars (\$63) commencing on July 1, 1983.

15 (c) The application fee for a clinical laboratory scientist's or
16 limited clinical laboratory scientist's license is thirty-eight dollars
17 (\$38) commencing on July 1, 1983.

18 (d) The application and annual renewal fee for a
19 cytotechnologist's license is fifty dollars (\$50) commencing on
20 January 1, 1991.

21 (e) The annual renewal fee for a clinical laboratory scientist's
22 or limited clinical laboratory scientist's license is twenty-five
23 dollars (\$25) commencing on July 1, 1983.

24 (f) A clinical laboratory applying for a license to perform tests
25 or examinations classified as of moderate or of high complexity
26 under CLIA and a clinical laboratory applying for certification
27 under subdivision (c) of Section 1223 shall pay an application fee
28 for that license or certification based on the number of tests it
29 performs or expects to perform in a year, as follows:

30 (1) Less than 2,001 tests: two hundred seventy dollars (\$270).

31 (2) Between 2,001 and 10,000, inclusive, tests: eight hundred
32 twenty dollars (\$820).

33 (3) Between 10,001 and 25,000, inclusive, tests: one thousand
34 three hundred fifteen dollars (\$1,315).

35 (4) Between 25,001 and 50,000, inclusive, tests: one thousand
36 five hundred eighty dollars (\$1,580).

37 (5) Between 50,001 and 75,000, inclusive, tests: one thousand
38 nine hundred sixty dollars (\$1,960).

39 (6) Between 75,001 and 100,000, inclusive, tests: two thousand
40 three hundred forty dollars (\$2,340).

1 (7) Between 100,001 and 500,000, inclusive, tests: two thousand
2 seven hundred forty dollars (\$2,740).

3 (8) Between 500,001 and 1,000,000, inclusive, tests: four
4 thousand nine hundred ten dollars (\$4,910).

5 (9) More than 1,000,000 tests: five thousand two hundred sixty
6 dollars (\$5,260) plus three hundred fifty dollars (\$350) for every
7 500,000 tests over 1,000,000, up to a maximum of 15,000,000
8 tests.

9 (g) A clinical laboratory performing tests or examinations
10 classified as of moderate or of high complexity under CLIA and
11 a clinical laboratory with a certificate issued under subdivision (c)
12 of Section 1223 shall pay an annual renewal fee based on the
13 number of tests it performed in the preceding calendar year, as
14 follows:

15 (1) Less than 2,001 tests: one hundred seventy dollars (\$170).

16 (2) Between 2,001 and 10,000, inclusive, tests: seven hundred
17 twenty dollars (\$720).

18 (3) Between 10,001 and 25,000, inclusive, tests: one thousand
19 one hundred fifteen dollars (\$1,115).

20 (4) Between 25,001 and 50,000, inclusive, tests: one thousand
21 three hundred eighty dollars (\$1,380).

22 (5) Between 50,001 and 75,000, inclusive, tests: one thousand
23 seven hundred sixty dollars (\$1,760).

24 (6) Between 75,001 and 100,000, inclusive, tests: two thousand
25 forty dollars (\$2,040).

26 (7) Between 100,001 and 500,000, inclusive, tests: two thousand
27 four hundred forty dollars (\$2,440).

28 (8) Between 500,001 and 1,000,000, inclusive, tests: four
29 thousand six hundred ten dollars (\$4,610).

30 (9) More than 1,000,000 tests per year: four thousand nine
31 hundred sixty dollars (\$4,960) plus three hundred fifty dollars
32 (\$350) for every 500,000 tests over 1,000,000, up to a maximum
33 of 15,000,000 tests.

34 (h) The application fee for a trainee's license is thirteen dollars
35 (\$13) commencing on July 1, 1983.

36 (i) The annual renewal fee for a trainee's license is eight dollars
37 (\$8) commencing on July 1, 1983.

38 (j) The application fee for a duplicate license is five dollars (\$5)
39 commencing on July 1, 1983.

1 (k) The personnel licensing delinquency fee is equal to the
2 annual renewal fee.

3 (l) The director may establish a fee for examinations required
4 under this chapter. The fee shall not exceed the total cost to the
5 department in conducting the examination.

6 (m) A clinical laboratory subject to registration under paragraph
7 (2) of subdivision (a) of Section 1265 and performing only those
8 clinical laboratory tests or examinations considered waived under
9 CLIA shall pay an annual fee of one hundred dollars (\$100). A
10 clinical laboratory subject to registration under paragraph (2) of
11 subdivision (a) of Section 1265 and performing only
12 provider-performed microscopy, as defined under CLIA, shall pay
13 an annual fee of one hundred fifty dollars (\$150). A clinical
14 laboratory performing both waived and provider-performed
15 microscopy shall pay an annual registration fee of one hundred
16 fifty dollars (\$150).

17 (n) The costs of the department in conducting a complaint
18 investigation, imposing sanctions, or conducting a hearing under
19 this chapter shall be paid by the clinical laboratory. The fee shall
20 be no greater than the fee the laboratory would pay under CLIA
21 for the same type of activities and shall not be payable if the
22 clinical laboratory would not be required to pay those fees under
23 CLIA.

24 (o) The state, a district, city, county, city and county, or other
25 political subdivision, or any public officer or body shall be subject
26 to the payment of fees established pursuant to this chapter or
27 regulations adopted thereunder.

28 (p) In addition to the payment of registration or licensure fees,
29 a clinical laboratory located outside the State of California shall
30 reimburse the department for travel and per diem to perform any
31 necessary onsite inspections at the clinical laboratory in order to
32 ensure compliance with this chapter.

33 (q) The department shall establish an application fee and a
34 renewal fee for a medical laboratory technician license, the total
35 fees collected not to exceed the costs of the department for the
36 implementation and operation of the program licensing and
37 regulating medical laboratory technicians pursuant to Section
38 1260.3.

39 (r) The costs of the department to conduct any reinspections to
40 ensure compliance of a laboratory applying for initial licensure

shall be paid by the laboratory. This additional cost for each visit shall be equal to the initial application fee and shall be paid by the laboratory prior to issuance of a license. The department shall not charge a reinspection fee if the reinspection is due to error or omission on the part of the department.

(s) A fee of twenty-five dollars (\$25) shall be assessed for approval of each additional location authorized by paragraph (2) of subdivision (d) of Section 1265.

(t) ~~Beginning in fiscal year 2013-14 On or before July 1, 2013,~~ the department shall report during the annual legislative budget hearing process ~~on the status of the department's work toward receiving an exemption from federal CLIA oversight, any deficiencies identified by the department, the extent to which the state oversight program meets or exceeds federal oversight standards~~ and the extent to which the federal Department of Health and Human Services is accepting exemption applications and the *potential* cost to the state for an exemption. ~~Reporting under this subdivision shall no longer be required when the state receives an exemption from the federal Department of Health and Human Services for federal CLIA oversight.~~

~~SEC. 6.~~

SEC. 5. Section 1300.2 is added to the Business and Professions Code, to read:

1300.2. Notwithstanding any other provision of this article, the total fees collected under this chapter shall not exceed the costs incurred by the department for licensing, certification, inspection, or other activities relating to the regulation of clinical laboratories and their personnel.

~~SEC. 7.~~

SEC. 6. Section 1301 of the Business and Professions Code is amended to read:

1301. (a) The annual renewal fee for a clinical laboratory license or registration set under this chapter shall be paid during the 30-day period before the expiration date of the license or registration. If the license or registration is not renewed before the expiration date, the licensee or registrant, as a condition precedent to renewal, shall pay a delinquency fee equal to 25 percent of the annual renewal fee for up to 60 days after the expiration date, in addition to the annual renewal fee in effect on the last preceding regular renewal date. Failure to pay the annual renewal fee in

1 advance during the time the license or registration remains in force
2 shall, ipso facto, work a forfeiture of the license or registration
3 after a period of 60 days from the expiration date of the license or
4 registration.

5 (b) (1) The department shall give written notice to all persons
6 licensed pursuant to Sections 1260, 1260.1, 1261, 1261.5, 1262,
7 1264, or 1270 30 days in advance of the regular renewal date that
8 a renewal fee has not been paid. In addition, the department shall
9 give written notice to licensed clinical laboratory bioanalysts or
10 doctoral degree specialists and clinical laboratory scientists or
11 limited clinical laboratory scientists by registered or certified mail
12 90 days in advance of the expiration of the fifth year that a renewal
13 fee has not been paid and if not paid before the expiration of the
14 fifth year of delinquency the licensee may be subject to
15 reexamination.

16 (2) If the renewal fee is not paid for five or more years, the
17 department may require an examination before reinstating the
18 license, except that no examination shall be required as a condition
19 for reinstatement if the original license was issued without an
20 examination. No examination shall be required for reinstatement
21 if the license was forfeited solely by reason of nonpayment of the
22 renewal fee if the nonpayment was for less than five years.

23 (3) If the license is not renewed within 60 days after its
24 expiration, the licensee, as a condition precedent to renewal, shall
25 pay the delinquency fee identified in subdivision (l) of Section
26 1300, in addition to the renewal fee in effect on the last preceding
27 regular renewal date. Payment of the delinquency fee will not be
28 necessary if within 60 days of the license expiration date the
29 licensee files with the department an application for inactive status.

30 ~~SEC. 8.— Section 1302 of the Business and Professions Code is~~
31 ~~amended to read:~~

32 ~~1302. (a) There is hereby established in the State Treasury,~~
33 ~~the Clinical Laboratory Improvement Fund.~~

34 ~~(b) All fees established under this chapter, Chapter 4~~
35 ~~(commencing with Section 1600) of Division 2 of the Health and~~
36 ~~Safety Code, and Article 5 (commencing with Section 101150) of~~
37 ~~Chapter 2 of Part 3 of Division 101 of the Health and Safety Code,~~
38 ~~shall be collected by and paid to the department, and shall be~~
39 ~~deposited by the department in the Clinical Laboratory~~
40 ~~Improvement Fund, along with any other moneys received by the~~

1 department for the purpose of licensing, certification, inspection,
2 proficiency testing, or other regulation of clinical laboratories;
3 public health laboratories, blood banks, or clinical or public health
4 laboratory personnel. Notwithstanding Section 16305.7 of the
5 Government Code, all interest earned on moneys deposited in the
6 fund shall be maintained in the fund.

7 ~~(c) Notwithstanding Section 13340 of the Government Code,~~
8 ~~moneys deposited in the Clinical Laboratory Improvement Fund~~
9 ~~are hereby continuously appropriated to the department to~~
10 ~~administer this chapter, Chapter 4 (commencing with Section 1600)~~
11 ~~of Division 2 of the Health and Safety Code, and Article 5~~
12 ~~(commencing with Section 101150) of Chapter 2 of Part 3 of~~
13 ~~Division 101 of the Health and Safety Code, and shall not be~~
14 ~~redirected for any other purpose. All fees collected pursuant to~~
15 ~~this chapter shall be expended to administer this chapter. All fees~~
16 ~~collected pursuant to Chapter 4 (commencing with Section 1600)~~
17 ~~of Division 2 of the Health and Safety Code shall be expended to~~
18 ~~administer that chapter. All fees collected pursuant to Article 5~~
19 ~~(commencing with Section 101150) of Chapter 2 of Part 3 of~~
20 ~~Division 101 of the Health and Safety Code shall be expended to~~
21 ~~administer that article.~~

22 *SEC. 7. Section 1302 of the Business and Professions Code is*
23 *amended to read:*

24 1302. (a) There is hereby established in the State Treasury,
25 the Clinical Laboratory Improvement Fund.

26 (b) All fees established under this chapter and Chapter 4
27 (commencing with Section 1600) of Division 2 of the Health and
28 Safety Code shall be collected by and paid to the department, and
29 shall be deposited by the department in the Clinical Laboratory
30 Improvement Fund, along with any other moneys received by the
31 department for the purpose of licensing, certification, inspection,
32 proficiency testing, or other regulation of clinical laboratories,
33 blood banks, or clinical laboratory personnel. *Notwithstanding*
34 *Section 16305.7 of the Government Code, all interest earned on*
35 *moneys deposited in the fund shall be maintained in the fund.*

36 (c) Moneys deposited in the Clinical Laboratory Improvement
37 Fund that are appropriated in the annual Budget Act, or any other
38 appropriation, for support of, or expenditure by, the state
39 department shall, upon appropriation, be expended by the state
40 department to administer this chapter and Chapter 4 (commencing

1 with Section 1600) of Division 2 of the Health and Safety Code.
2 All fees collected pursuant to this chapter shall, upon appropriation,
3 be expended to administer this chapter *and shall not be redirected*
4 *for any other purpose*. All fees collected pursuant to Chapter 4
5 (commencing with Section 1600) of Division 2 of the Health and
6 Safety Code shall, upon appropriation, be expended to administer
7 that chapter *and shall not be redirected for any other purpose*.

8 ~~SEC. 9.~~

9 *SEC. 8.* This act is an urgency statute necessary for the
10 immediate preservation of the public peace, health, or safety within
11 the meaning of Article IV of the Constitution and shall go into
12 immediate effect. The facts constituting the necessity are:

13 In order to protect the public health by providing strong clinical
14 laboratory oversight as soon as possible, it is necessary that this
15 act take effect immediately.