Follow-Up—California Department of Public Health

Laboratory Field Services Is Unable to Oversee Clinical Laboratories Effectively, but a Feasible Alternative Exists

Report 2015-507
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September 10, 2015

The Governor of California  
President pro Tempore of the Senate  
Speaker of the Assembly  
State Capitol  
Sacramento, California 95814

Dear Governor and Legislative Leaders:

This report presents the results of a follow-up audit the California State Auditor conducted concerning the efforts by Laboratory Field Services (Laboratory Services)—within the California Department of Public Health (Public Health)—to implement recommendations from an audit report that we issued in September 2008. The report titled Department of Public Health: Laboratory Field Services’ Lack of Clinical Laboratory Oversight Places the Public at Risk, Report 2007-040, examined Laboratory Services’ ability to oversee clinical laboratories (labs) that analyze human specimens such as blood, tissue, and urine so that medical professionals can make diagnoses and prescribe treatment.

In this follow-up audit, we found that Laboratory Services is still not performing the oversight activities with which it is entrusted and its management of the program remains inadequate. Laboratory Services has not fully implemented many of the recommendations from our September 2008 audit report. Laboratory Services still only inspects approximately half of California labs and it does not have a process to ensure that it is aware, in a timely manner, when out-of-state labs that are licensed in California fail required proficiency testing. Laboratory Services also continues to not investigate all the complaints it receives and has issued only a small number of sanctions in the past seven years even though it is responsible for overseeing more than 22,100 labs. Moreover, we found that Laboratory Services made an unauthorized fee increase in January 2014 that has resulted in labs overpaying it more than $1 million in fees, and since 2008 it has collected more than $12 million in lab fees that it has not spent. Finally, Laboratory Services has missed opportunities to more effectively use its limited personnel by partnering with other organizations that could help it meet its workload obligations under state law.

Since the problems that have plagued Laboratory Services have persisted since our last audit, we believe the State’s consumers have, in effect, been relying on federal oversight the Centers for Medicare and Medicaid Services (CMS) provides through its administration of the Clinical Laboratory Improvement Amendments of 1988 (CLIA). In fact, we believe Laboratory Services’ oversight of lab facilities largely duplicates federal oversight with no meaningful benefit to consumers. State law and CLIA are nearly equivalent in their mandates, and the oversight required is redundant: Both Laboratory Services and CMS collect fees from labs to perform inspections, monitor proficiency testing, investigate complaints, and issue sanctions. The Legislature should repeal state law requiring that lab facilities be licensed by the State, thus reducing the regulatory and financial burden on lab facilities while continuing to enforce the State’s requirements for laboratory personnel.

Respectfully submitted,

ELAINE M. HOWLE, CPA  
State Auditor
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California Department of Public Health 47
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Summary

Results in Brief

Laboratory Field Services (Laboratory Services) within the California Department of Public Health (Public Health) is responsible for overseeing clinical laboratories (labs) that analyze human specimens such as blood, tissue, and urine. Medical professionals use these analyses to make diagnoses and prescribe treatment. Laboratory Services’ oversight responsibilities cover both labs located within California and labs located outside of the State that test specimens originating from within California. The State currently has licensed approximately 2,800 labs and registered approximately 19,300 labs; the complexity of the tests the labs perform dictates whether they require licensing or registration. Laboratory Services’ oversight responsibilities include inspecting licensed labs once every two years and periodically verifying the accuracy and reliability of their tests through a process called proficiency testing. It must also investigate complaints against both licensed and registered labs and may issue sanctions when it finds that a lab is out of compliance with state laws or regulations. All licensed labs must pay Laboratory Services an annual fee based on the volume of tests they perform, while registered labs must pay an annual flat fee.

In this follow-up audit, we found that Laboratory Services is still not performing the oversight activities with which it has been entrusted and that its management of its responsibilities is inadequate. Specifically, it has not implemented many of the recommendations from our September 2008 audit report titled Department of Public Health: Laboratory Field Services’ Lack of Clinical Laboratory Oversight Places the Public at Risk, Report 2007-040 (2008 audit). For example, it still only inspects about half of California labs, and it has not established a process to ensure that it becomes aware, in a timely manner, when out-of-state labs that are licensed in California fail required proficiency testing. Further, it does not yet investigate all complaints against labs and has issued only a small number of lab sanctions in the past seven years despite the number of labs it oversees. Additionally, we found that Laboratory Services made an unauthorized fee increase in January 2014 that resulted in labs overpaying it more than $1 million, and since 2008 it has collected more than $12 million in lab fees that it has not spent. Finally, its management has missed opportunities to more effectively use its limited personnel by partnering with other organizations that could help it meet its workload obligations under state law. Under state law, Laboratory Services can approve private nonprofit accreditation organizations to conduct oversight functions—including performing inspections.

Audit Highlights . . .

Our follow-up audit of Laboratory Field Services’ (Laboratory Services) progress in addressing issues we raised in our September 2008 report revealed the following:

» Laboratory Services has not implemented many of the recommendations from our prior audit report.

» Laboratory Services is still not performing oversight activities of clinical laboratories (labs) required by law and its management of its responsibilities is inadequate.

• It only inspects about half of California labs, and has not established a process to ensure that it becomes promptly aware when California licensed out-of-state labs fail required proficiency testing.

• It does not investigate all complaints against labs and has issued only a small number of lab sanctions in the past seven years despite the number of labs it oversees.

• It made an unauthorized fee increase in January 2014, resulting in labs overpaying it more than $1 million and contributing to a $12 million fund balance it has not spent.

• Management has missed opportunities to partner with nonprofit accreditation organizations to conduct oversight functions, inspections, and monitor proficiency testing.

» To reduce the regulatory and financial burden on lab facilities, state law requiring that lab facilities be licensed by the State should be repealed.
and monitoring proficiency testing—in lieu of its direct oversight. However, Laboratory Services has not taken full advantage of this opportunity.

Because the problems that have plagued Laboratory Services have persisted since our last audit, we believe the State’s consumers have, in effect, been relying on the federal oversight that the federal Centers for Medicare and Medicaid Services (CMS) provides through its administration of the Clinical Laboratory Improvement Amendments of 1988 (CLIA). In fact, we believe Laboratory Services’ workload largely duplicates federal oversight with no meaningful benefit to consumers. State law and CLIA are nearly equivalent in their mandates, and the oversight each requires is redundant: Both Laboratory Services and CMS collect fees from labs to perform inspections, monitor proficiency testing, investigate complaints, and issue sanctions. Eliminating the portion of Laboratory Services that we have found to be exceedingly deficient for many years would end the duplicate oversight and the duplicate fees. CMS has processes in place to ensure that it effectively administers CLIA; therefore, relying on CLIA would keep the public health benefits of lab monitoring intact, as they are today, while reducing the regulatory burden on California’s clinical labs.

**Recommendations**

To eliminate the State’s redundant and ineffective oversight of labs and to ensure labs do not pay unnecessary or duplicative fees, the Legislature should do the following:

- Repeal existing state law requiring that labs be licensed or registered by Laboratory Services and that Laboratory Services perform oversight of these labs. Instead, the State should rely on the oversight CMS provides.

- Repeal existing state law requiring labs to pay fees for state-issued licenses or registrations.

While the Legislature considers eliminating the requirements that labs receive state-issued licenses or registrations and that Laboratory Services oversee these labs, Laboratory Services should develop a corrective action plan by December 31, 2015. This plan should identify the individuals responsible for ensuring Laboratory Services takes corrective actions, the resources needed to carry out those corrective actions, and the expected time frame.
for their successful implementation. The corrective action plan should address Laboratory Services’ plans for implementing the recommendations from our 2008 audit, including the following:

- Inspecting licensed labs every two years and ensuring it identifies in a timely manner all labs that fail proficiency testing. In addition, it should improve its complaints policy and procedures and dedicate an appropriate number of staff to its sanctioning efforts.

- Developing a process to assess the budget act annually and to adjust its fees accordingly. It should also maximize the opportunity to partner with accreditation organizations by developing an accreditation organization program.

Agency Comments

Public Health responded that it concurred with the recommendations and outlined a number of steps it will take to implement them.
Introduction

Background

Clinical laboratories (labs) analyze human specimens such as blood, tissue, and urine so that medical professionals can make diagnoses and prescribe treatment. As a part of the California Department of Public Health (Public Health) and under the direction of the Office of the State Public Health Laboratory Director, Laboratory Field Services (Laboratory Services) is responsible for licensing, registering, and overseeing labs. As of July 2015 Laboratory Services reported it was responsible for overseeing roughly 22,100 labs.

According to state law, the complexity of the tests that labs perform determines whether they must obtain licenses or registrations, as summarized in the text box. Of the approximately 22,100 clinical labs Laboratory Services was responsible for overseeing as of July 2015, about 2,800 were licensed and about 19,300 were registered. A license or registration is valid for one year, thus requiring annual renewal for the lab to continue operating.

A lab seeking to obtain or renew a license or registration must pay a fee to Laboratory Services. Although registration fees are a set amount, each lab’s license fee is based on the volume of tests it conducts. Laboratory Services deposits the fees and other money it collects into the Clinical Laboratory Improvement Fund. The law states that the total fees Laboratory Services collects shall not exceed its costs for licensing, certifying, and inspecting labs, as well as performing other activities relating to the regulation of labs and lab personnel. For fiscal year 2013–14, Laboratory Services reported more than $6.5 million in fee revenue from licensed and registered labs.

At times, a medical professional located in California will send a specimen to a lab in another state or another country for analysis; these labs are referred to as out-of-state labs. State law requires that the receiving lab hold a license or registration from Laboratory Services. Further, out-of-state labs that Laboratory Services licenses rather than registers are subject to its periodic oversight, as described below.

### Registration and Licensure Requirements for Clinical Laboratories

- Clinical laboratories (labs) requiring licensure perform tests of moderate to high complexity, such as testing for hepatitis or certain sexually transmitted diseases by DNA probe.
- Labs requiring registration perform simpler tests, with less chance of error or risk, such as prepackaged manufactured tests.

Sources: California Business and Professions Code, Section 1265, and Laboratory Field Services’ documents.

State-Mandated Responsibilities for Lab Oversight

The State has overseen labs since 1926 and has licensed labs since the 1950s. State law currently requires Laboratory Services to oversee labs by inspecting them, monitoring their proficiency...
testing, annually renewing their licenses and registrations, receiving and investigating complaints against them, and sanctioning those that violate laws or regulations. Laboratory Services must engage in two periodic oversight functions: conducting regular inspections and monitoring proficiency testing. According to state law, Laboratory Services must inspect each licensed lab every two years, notify the lab of any deficiencies the inspection reveals, and work with the lab to correct the deficiencies. Registered labs are not subject to routine inspections every two years under state law, but Laboratory Services may inspect them as part of complaint investigations.

Laboratory Services has offices in Richmond and Los Angeles. It divides the licensing, registration, and oversight functions it is mandated to perform between the two locations. Figure 1 is a

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**Proficiency Testing Process for Licensed Clinical Laboratories**

**What Is Proficiency Testing?**
Proficiency testing is a process clinical laboratories (labs) use to verify the accuracy and reliability of their tests.

**How Does Proficiency Testing Work?**
A provider distributes a specimen to a lab, which must evaluate the specimen and then submit the results to the provider. The provider has a target value for the specimen, and on receiving the lab's assessment, the provider compares the lab's results with its target value to determine if the lab's evaluation was accurate.

**How Often Must Labs Test?**
In general, labs must engage in proficiency testing at least three times a year.

**What Is a Testing Failure?**
Participation is unsuccessful if the lab does not achieve a minimum score on either two consecutive tests or two out of three consecutive tests.

Sources: California Business and Professions Code, Section 1220, and Title 42, Code of Federal Regulations, Part 493.

The second type of periodic oversight Laboratory Services must perform is monitoring proficiency-testing results. Proficiency testing provides an external evaluation of the accuracy of the labs' test results. Licensed labs must participate because they perform complex tests; however, registered labs—which perform simple tests—are not required to participate in proficiency testing. The text box describes the proficiency testing process. Laboratory Services' policy generally calls for it to receive and review each lab's proficiency-testing results at least three times a year and identify any instances of unsatisfactory performance. In those instances, according to its policy, Laboratory Services must notify the lab and require a plan of corrective action. If the planned corrective action is not acceptable or the lab's test results do not improve, Laboratory Services can bar the lab from providing those test services.

Laboratory Services' other oversight responsibilities include investigating complaints and issuing sanctions. State law requires Laboratory Services to investigate complaints it receives about labs and authorizes it to inspect labs as part of its complaint investigations. Further, when labs do not adhere to state law and regulations, Laboratory Services has the authority to issue sanctions that can include monetary penalties, plans of correction, and license or registration revocation. If Laboratory Services revokes a lab's license or registration, the lab's owner and operator are automatically barred from owning or operating a lab for two years.

Laboratory Services has offices in Richmond and Los Angeles. It divides the licensing, registration, and oversight functions it is mandated to perform between the two locations. Figure 1 is a
partial depiction of Laboratory Services’ organizational structure. As the figure shows, two of Laboratory Services’ sections perform functions related to the state mandates for labs. One section, located in Los Angeles, oversees federal lab requirements, as described below.

**Figure 1**
Partial Depiction of Laboratory Field Services’ Organizational Structure as of February 2015

Sources: Laboratory Field Services’ organization chart dated February 18, 2015, and the California State Auditor’s analysis of functions assigned to each section.

**Federal Oversight of Labs**

In addition to meeting state requirements, all the labs that Laboratory Services licenses or registers must also follow federal regulations. The Clinical Laboratory Improvement Amendments of 1988 (CLIA) is a federal law enacted to ensure the accuracy and reliability of lab testing. This law extended federal regulation for the first time to all labs in the nation that perform tests on human specimens for medical diagnosis, treatment, or health assessment. The federal Centers for Medicare and Medicaid Services (CMS) has primary responsibility under CLIA for regulating approximately 250,000 labs nationwide as of November 2014. CMS meets this responsibility in part by contracting with state agencies across the country to monitor and enforce compliance with CLIA. By law, activities to enforce CLIA
requirements must be self-funded. With few exceptions, labs must apply for a CLIA certificate and pay a biennial fee to cover the cost of inspections and other regulatory activities.

CLIA groups labs into two categories—those performing simple tests, such as urine dipstick tests and finger-stick blood tests, and those performing moderately complex to highly complex tests (complex tests). A lab’s category dictates the federal oversight to which it is subject. CLIA exempts labs from virtually all federal rules if they perform only simple tests in strict compliance with the manufacturers’ instructions. However, as Figure 2 shows, labs that perform complex tests differ from those performing simple tests in two ways: They are subject to ongoing oversight in the form of biennial inspections and proficiency testing, and they can choose their oversight body.

**Figure 2**
Clinical Laboratory Improvement Amendments of 1988—Requirements and Oversight

![Diagram showing the requirements and oversight for simple and complex tests.](image-url)


* CMS has primary responsibility for administering the Clinical Laboratory Improvement Amendments of 1988, which it accomplishes through contracts with state agents.

† Accreditation organizations chosen to oversee a licensed clinical laboratory must be approved by CMS.
Labs that perform complex tests can choose to be monitored directly by CMS through the state agencies with which it contracts or they can voluntarily apply for accreditation from private, nonprofit, CMS-approved accreditation organizations. Although CMS has primary responsibility for administering CLIA, it contracts with state governments to provide the federally required oversight of nonaccredited labs, as we previously mentioned. We refer to the state agencies that provide federal CLIA oversight as state agents. California, through Public Health, has entered into an agreement to act as CMS's state agent to oversee nonaccredited labs within the State. Therefore, in addition to its responsibilities related to state clinical lab law, Laboratory Services performs CLIA-related duties as the state agent for CMS. As Figure 1 on page 7 shows, a specific section of Laboratory Services based in its Los Angeles office acts as the CMS state agent (CLIA section). We use the term Laboratory Services throughout this report to refer to the sections of Laboratory Services that perform its state-mandated responsibilities, not its federal responsibilities per its agreement with CMS.

A lab that performs complex tests that seeks accreditation from an accreditation organization is directly overseen by that organization. If an accredited lab complies with its accreditation organization's requirements, CMS deems it as meeting all applicable CLIA requirements. A lab seeking accreditation must also apply to and pay CMS for a CLIA certificate in addition to any fees or registration requirements imposed by its accreditation organization. Figure 3 on the following page illustrates the various entities that oversee labs and identifies whether those entities monitor compliance with state or federal lab requirements.

**Scope and Methodology**

California Government Code, Section 8546.1(d), authorizes the California State Auditor (state auditor) to conduct follow-up audit work on statutorily mandated or legislatively requested financial and performance audits. In September 2008 the state auditor published a report titled *Department of Public Health: Laboratory Field Services’ Lack of Clinical Laboratory Oversight Places the Public at Risk*, Report 2007-040. In March 2015 the state auditor initiated a follow-up audit to evaluate whether Laboratory Services had improved its oversight of labs. Table 1 beginning on page 11 lists those recommendations from the 2008 report on which we followed up and our methods for assessing their implementation status.
**Figure 3**
Clinical Laboratory Oversight in California for Laboratories That Perform Complex Tests

**Source**: California State Auditor's analysis of state and federal oversight structures of labs.

CLIA = Clinical Laboratory Improvement Amendments of 1988.

* Labs can choose oversight by either private nonprofit accreditation organizations approved by CMS or through direct CMS oversight via the CLIA section.

† CMS contracts with states to provide federally required oversight of nonaccredited labs. Laboratory Services' CLIA section is responsible for ensuring that California labs comply with CLIA requirements. It also reviews accreditation organizations' performance on behalf of CMS.

‡ Laboratory Services' facility licensing sections include offices in Richmond and Los Angeles that license and oversee labs.
Table 1
Methods Used to Review the Current Status of Recommendations From the September 2008 Audit Report 2007-040

<table>
<thead>
<tr>
<th>RECOMMENDATION</th>
<th>METHOD</th>
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<tbody>
<tr>
<td>1 Laboratory Field Services (Laboratory Services) should perform all its mandated oversight responsibilities for clinical laboratories (labs) subject to its jurisdiction operating within and outside California, including but not limited to the following: • Inspecting licensed labs every two years. • Monitoring proficiency-testing results. • Sanctioning labs as appropriate. • Reviewing and investigating complaints and ensuring necessary resolution.</td>
<td>• Interviewed key Laboratory Services officials about its inspections practices and reviewed relevant laws. • Obtained lab inspection data and analyzed them to determine whether Laboratory Services met its mandate to inspect licensed labs every two years. • Performed file reviews to ascertain the reliability of the lab inspection data. We noted data reliability concerns with the data but determined our concerns would not change our conclusions. • Identified state oversight mandates that duplicate federal oversight requirements under the Clinical Laboratory Improvement Amendments of 1988. • Evaluated Laboratory Services' proficiency testing, sanctions, and complaint investigations as described below.</td>
</tr>
<tr>
<td>2 Laboratory Services should adopt and implement proficiency-testing policies and procedures for staff to do the following: • Promptly review labs' proficiency-testing results and notify labs that fail. • Follow specified timelines for responding to labs' attempts to correct proficiency-testing failures and for sanctioning labs that do not comply. • Monitor the proficiency-testing results of out-of-state labs. • Verify labs' enrollment in proficiency testing, and ensure that it receives proficiency-testing scores from all enrolled labs.</td>
<td>• Interviewed key Laboratory Services officials about its practices for monitoring proficiency-testing results and reviewed laws, policies, and procedures. • Reviewed labs' proficiency-testing results and examined 10 results that were deficient to determine what actions Laboratory Services had taken to ensure the labs corrected their deficiencies.</td>
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<tr>
<td>3 To update its regulations, Laboratory Services should review its clinical lab regulations and repeal or revise them as necessary. As part of its efforts to revise its regulations, Laboratory Services should ensure that they include requirements such as the time frames it wants to impose on the lab community.</td>
<td>• Interviewed key officials from Laboratory Services, the Office of the State Public Health Laboratory Director, and the California Department of Public Health (Public Health) about regulations development. • Obtained and reviewed Public Health's regulations tracking logs to assess which regulations it has planned for development and the associated timelines.</td>
</tr>
<tr>
<td>4 To strengthen its complaints process, Laboratory Services should identify necessary controls and incorporate them into its complaints policies. The necessary controls include, but are not limited to, receiving, logging, tracking, and prioritizing complaints, as well as ensuring that substantiated allegations are corrected. In addition, Laboratory Services should develop and implement corresponding procedures for each control. Further, Laboratory Services should establish procedures to ensure that it promptly forwards complaints for which it lacks jurisdiction to the entity that has jurisdiction.</td>
<td>• Interviewed key Laboratory Services officials regarding its complaints processes and reviewed its complaints policies and procedures. • Obtained complaints data and analyzed them to determine how many complaints Laboratory Services received from September 2008 through May 2015. • Performed file reviews to ascertain the reliability of the complaints data. We noted data reliability concerns with the data but determined our concerns would not change our conclusions. • Determined what actions Laboratory Services took to follow up on complaints by reviewing five open complaints. For five complaints that resulted in corrective action plans, we determined what Laboratory Services did to ensure the labs complied with the plans.</td>
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continued on next page...
## RECOMMENDATION

5 To strengthen its sanctioning efforts, Laboratory Services should do the following:
- Maximize its opportunities to impose sanctions.
- Appropriately justify and document the amounts of the civil money penalties it imposes.
- Ensure that it always collects the penalties it imposes.
- Follow up to ensure that labs take corrective action.
- Ensure that when it sanctions a lab, it notifies other appropriate agencies as necessary.

### METHOD
- Interviewed key Laboratory Services officials regarding its sanction policy and procedures, the sanctions it has issued, and its staffing for sanction activities.
- Obtained sanctions data and analyzed them to determine how many sanctions Laboratory Services has issued since 2008. We noted data reliability concerns with the data but determined our concerns would not change our conclusions. We reviewed hardcopy lab files.
- Compared sanctions data to official accounting records from Public Health and the California State Controller’s Office.

6 Public Health, in conjunction with Laboratory Services, should ensure that Laboratory Services has sufficient resources to meet all its oversight responsibilities.

### METHOD
- Interviewed key officials from Laboratory Services and the Office of the State Public Health Laboratory Director about staffing, succession planning, and planned reorganization.
- Obtained and reviewed supporting documentation.

7 Laboratory Services should work with its Information Technology Services Division and other appropriate parties to ensure that its data systems support its needs. If Laboratory Services continues to use its internally developed databases, it should ensure that it develops and implements appropriate system controls.

### METHOD
- Interviewed key officials from Laboratory Services, Public Health, and the Office of the State Public Health Laboratory Director to determine the updates that Laboratory Services has made to its existing information systems and its plans, if any, to replace those systems.
- Reviewed change logs to identify any changes Laboratory Services made to relevant information technology systems.

8 To demonstrate that it has used existing resources strategically and has maximized their utility to the extent possible, Laboratory Services should identify and explore opportunities to leverage existing processes and procedures. These opportunities should include, but not be limited to, exercising clinical lab oversight when it renews licenses and registrations, developing a process to share state concerns identified during federal inspections, and using accreditation organizations and contracts to divide its responsibilities for inspections every two years.

### METHOD
- Interviewed key officials from Laboratory Services about the steps it has taken to implement Senate Bill 744 (Chapter 201, Statutes of 2009).
- Reviewed relevant laws.
- Determined the number of accreditation organizations that applied to Laboratory Services, the number of accreditation organizations Laboratory Services approved and the time frames for approval.

9 Laboratory Services should work with Public Health’s budget section and other appropriate parties to ensure that it adjusts fees in accordance with the annual budget act.

### METHOD
- Interviewed key Laboratory Services officials and analyzed relevant laws and budget acts.
- Reviewed Laboratory Services’ financial statements and compared them to the California State Controller’s Office’s records.
- Compared Laboratory Services’ fee adjustments from fiscal years 2009–10 through 2014–15 to the fee adjustments required by the annual budget acts and identified the differences.
- Compared Laboratory Services’ license and registration revenue it collected for fiscal years 2008–09 through 2013–14 to its expenditures to determine whether it collected fees in excess of its operating needs.

Sources: Recommendations made in the report by the California State Auditor titled Department of Public Health: Laboratory Field Services’ Lack of Clinical Laboratory Oversight Places the Public at Risk, Report 2007-040, September 2008, and information and documentation identified in the table column titled Method.

### Assessment of Data Reliability

The U.S. Government Accountability Office, whose standards we are statutorily required to follow, requires us to assess the sufficiency and appropriateness of computer-processed information that is used to support our findings, conclusions, or recommendations. In our 2008 audit we found that Laboratory Services’ information technology systems—Health Applications Licensing system and four Microsoft Access databases, which
contained data for facility licenses, registrations, and certain oversight functions—did not adequately support Laboratory Services’ oversight activities or lacked the safeguards necessary to ensure accurate and complete information. Because of these known limitations, and because Laboratory Services had not fully implemented our 2008 recommendation to improve its information technology systems before the start of this follow-up audit, we did not conduct a data reliability assessment on Laboratory Services’ various management data. Therefore, Laboratory Services’ computer-processed information is of undetermined reliability for the purpose of this audit. Although this determination may affect the precision of the numbers we present, we believe we have gathered sufficient evidence in total to support our findings, conclusions, and recommendations.
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Audit Results

Laboratory Services Is Still Failing to Meet Its State Mandate to Oversee Clinical Laboratories

Over the last seven years, the California Department of Public Health’s (Public Health) Laboratory Field Services (Laboratory Services) has consistently failed to adequately oversee clinical laboratories (labs) as state law requires. In our September 2008 audit report titled Department of Public Health: Laboratory Field Services’ Lack of Clinical Laboratory Oversight Places the Public at Risk, Report 2007-040 (2008 audit), we found that Laboratory Services was not sufficiently inspecting labs, monitoring proficiency testing, investigating complaints, or issuing sanctions. Similarly, in this follow-up audit, we found that Laboratory Services has not inspected about half of the labs requiring such review under state law, and it continues to inconsistently monitor the results of proficiency testing for out-of-state labs. Additionally, Laboratory Services still struggles to investigate complaints promptly and issued only a small number of facility sanctions in the last seven years, although it oversees roughly 22,100 licensed and registered labs. Thus, it has not performed the oversight activities with which the State has entrusted it, as summarized in Table 2.

Table 2
Laboratory Field Services’ Implementation of the California State Auditor’s 2008 Recommendations Related to Its Oversight Responsibilities

<table>
<thead>
<tr>
<th>OVERSIGHT RESPONSIBILITY</th>
<th>2008 FINDING</th>
<th>RECOMMENDATION</th>
<th>RECOMMENDATION’S CURRENT STATUS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inspections</td>
<td>Laboratory Field Services (Laboratory Services) had not conducted any biennial inspections of licensed clinical laboratories (labs).</td>
<td>Laboratory Services should inspect licensed labs every two years.</td>
<td>Partially implemented.</td>
</tr>
<tr>
<td>Proficiency Testing</td>
<td>Laboratory Services did not consistently monitor labs. Its policy and procedures were inadequate, specifically with respect to monitoring out-of-state labs and reviewing proficiency-testing results in a timely manner. Laboratory Services did not identify or follow up on multiple deficiencies at labs.</td>
<td>Laboratory Services should adopt and implement policies and procedures for staff to promptly review proficiency-testing results and notify labs that fail. It should follow specified timelines for responding to labs’ attempts to correct failures and sanction labs that do not comply. Further, Laboratory Services should monitor results of out-of-state labs’ proficiency testing.</td>
<td>Partially implemented.</td>
</tr>
<tr>
<td>Complaints</td>
<td>Laboratory Services’ policy and procedures lacked sufficient safeguards to ensure that staff promptly logged and investigated complaints and ensured that labs correct substantiated allegations. It also closed some complaints without investigation.</td>
<td>Laboratory Services should update its policies and procedures to add safeguards over receiving, logging, tracking, and prioritizing complaints as well as ensuring that labs correct all substantiated allegations.</td>
<td>Partially implemented.</td>
</tr>
<tr>
<td>Sanctions</td>
<td>Laboratory Services did not always have staff dedicated to sanctioning efforts, lacked management data, and could not demonstrate that it collected the civil money penalties it imposed.</td>
<td>Laboratory Services should sanction labs as appropriate and strengthen its sanctioning efforts by justifying, documenting, and collecting the civil money penalties it imposes.</td>
<td>No action taken.</td>
</tr>
</tbody>
</table>

Sources: California State Auditor’s (state auditor) Report 2007-040 and the state auditor’s analysis of Laboratory Services’ corrective action.
Laboratory Services Is Not Inspecting Labs as State Law Requires

Laboratory Services is still not performing inspections of clinical labs as state law requires. According to state law, Laboratory Services must inspect all licensed labs biennially, or no less than once every two years. However, similar to our finding in 2008, Laboratory Services is not meeting its mandate, only inspecting about half of the labs it is required to inspect.

Over the last two years, Laboratory Services has not performed biennial inspections on a significant number of labs. In fact, it inspected only about half of the labs requiring biennial inspections each year.

Over the last two years, Laboratory Services has not performed biennial inspections on a significant number of labs. Just over 2,800 labs require biennial inspections; thus, Laboratory Services must inspect about 1,400 labs each year. However, Figure 4 shows the percentage of licensed labs that Laboratory Services inspected in 2013 and 2014, demonstrating that it inspected only about half of the labs requiring biennial inspections in each year. For workload purposes, Laboratory Services divides its biennial inspection responsibilities into three lab segments: out-of-state labs, in-state accredited labs, and in-state nonaccredited labs. Laboratory Services maintains responsibility for inspecting out-of-state and accredited labs but relies on its Clinical Laboratory Improvement Amendments of 1988 Section (CLIA section)—the unit that performs federal reviews on behalf of the federal Centers for Medicare and Medicaid Services (CMS)—to inspect in-state nonaccredited labs. Because state oversight requirements generally mirror CLIA, Laboratory Services views an inspection that its CLIA section performs as comparable.

Laboratory Services is far from meeting its obligation of inspecting all out-of-state and accredited labs once every two years. Its inspection rate for out-of-state labs—the smallest lab segment in its purview—is particularly problematic. State law has required out-of-state biennial inspections since 1996; however, the former chief of Laboratory Services reported that her staff did not begin performing out-of-state inspections until November 2014. The Los Angeles office performs out-of-state lab inspections and Figure 4, which we developed primarily from that office’s biennial inspection data, shows that Laboratory Services’ Los Angeles office did not inspect any out-of-state labs in 2013 and only five such labs in 2014. As a result, Laboratory Services failed to inspect at least 93 percent of out-of-state labs requiring biennial inspections in 2014. Going forward, Laboratory Services’ Los Angeles office plans to inspect two out-of-state labs each month. However, that plan will not reach the necessary number of lab inspections; instead, it will leave uninspected more than 60 percent of the 70 out-of-state labs requiring inspections.
Figure 4
Estimated Percentage of Clinical Laboratories That Received Either State or Federal Biennial Inspections in 2013 and 2014

Sources: California State Auditor’s analysis of Laboratory Field Services’ (Laboratory Services) inspection tracking logs and reports generated by the federal Centers for Medicare and Medicaid Services (CMS). Unaudited.

* Laboratory Services’ list of out-of-state labs for inspection numbered roughly 140 labs, which did not reconcile with its roughly 460 out-of-state licensed labs. Nevertheless, the number of out-of-state labs inspected is so low that the differences between the lists do not alter our conclusion.

† Laboratory Services’ CLIA section—the unit that enforces federal law titled the Clinical Laboratory Improvement Amendments of 1988 (CLIA) and performs federal reviews on behalf of CMS—did not meet its inspection mandate for biennial inspections in federal fiscal year 2014. The CMS performance review of that year recognized that the CLIA section operated with a shortage of inspectors during that year.

‡ When CLIA section examiners perform inspections related to federal oversight requirements, they also complete a state law checklist that allows those inspections to count towards the State’s mandate of inspecting labs biennially.

§ These data are estimated because we converted CMS data from federal fiscal years to calendar years, which required using averages.
State oversight of accredited labs is also lacking: Laboratory Services inspected less than 20 percent of these labs in the last two years, as shown in Figure 4. Laboratory Services divides inspections of accredited labs between its two offices and has assigned inspections for Northern California labs to one staff person in the Richmond office who performed only eight biennial inspections during 2013 and 2014. This staff person's logs show that he spent most of his time inspecting labs seeking state licensure as opposed to performing the recurring biennial inspections of currently licensed labs. Laboratory Services' data show that it has currently assigned more than 250 clinical labs to its Richmond office. Assuming that roughly 125 of these accredited labs must be inspected each year, the single staff person in Richmond performing between three and five biennial inspections annually makes a negligible impact on the required workload.

Although Laboratory Services' Los Angeles office has assigned inspections to a number of staff who perform significantly more biennial inspections of accredited labs than the single staff person in the Richmond office, it still falls far short of meeting its required workload. Data that Laboratory Services provided indicate that it has more than 600 clinical labs in Southern California that require recurring biennial inspections, translating to a need for it to perform roughly 300 biennial inspections each year. However, in 2013 and 2014, the Los Angeles staff only performed 87 and 108 biennial inspections, respectively, well under half of the required workload.

To meet its state mandate, Laboratory Services relies extensively on inspections that staff in its CLIA section perform. The CLIA section has seven state staff who perform federal-based reviews on behalf of CMS. The CLIA section is shown in Figure 1 on page 7 and is described in the Introduction. By counting CLIA inspections toward its requirement to perform recurring state biennial inspections, Laboratory Services has increased the number of labs inspected for state standards to around 50 percent, as Figure 4 on page 17 shows. In response to one of our previous recommendations to improve efficiency, Laboratory Services began including CLIA inspections in its counts of completed biennial inspections as early as November 2008. When CLIA section staff perform federal lab inspections, they complete a one-page state law checklist that Laboratory Services developed to determine whether the lab is also compliant with state standards. The

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1 Throughout the audit, we noted concerns and inconsistencies with the reliability of Laboratory Services' data. We discuss some of these concerns later in the report. Nevertheless, we determined that the questionable data would not change our conclusions regarding Laboratory Services' inability to meet its inspections workload. Moreover, in an internal memo, Laboratory Services acknowledges that it was not meeting that workload.
state law checklist includes 15 criteria—some of which relate to requirements for lab personnel rather than lab facilities—that differ between California and federal laws and regulations. The CLIA section inspected 694 labs in 2013 and 577 labs in 2014. Without the inspections the CLIA section performed, the percentage of all clinical labs that received state biennial inspections would be significantly lower.

Laboratory Services claims that staffing is the main reason it has not inspected labs as required. The acting facility section chief in the Los Angeles office stated that she suggested hiring more staff as well as increasing the number of biennial inspections each staff member must perform every month in order to increase the total number of inspected labs. However, as we describe in a later section of the report, Laboratory Services has had both the funding and the opportunity to hire more staff since we completed the 2008 audit, but it has not done so.

Additionally, the former Laboratory Services chief stated that Laboratory Services did not perform out-of-state inspections before November 2014 because of the governor’s restriction on out-of-state travel. In April 2011 Governor Brown issued an executive order requiring that his office approve all out-of-state travel. However, we believe based on the criteria included in the executive order that Laboratory Services likely would have qualified for an out-of-state travel exemption. For example, the executive order stated that no out-of-state travel would be permitted unless it was at no cost to the State or was “mission critical,” meaning that, for instance, it pertained to a department’s enforcement responsibilities or to a function required by statute. Because state law mandates the biennial inspections, we believe that travel related to them would have qualified as mission critical. Moreover, state law requires that out-of-state labs reimburse Laboratory Services for travel and per diem costs to perform any necessary on-site inspections; thus, the inspections would have incurred no travel costs to the State. Although Laboratory Services management believed it submitted out-of-state exemption requests, Laboratory Services and Public Health officials were unable to provide documentation of these requests. Further, the former Laboratory Services chief stated that as of May 2015 Laboratory Services had not sought reimbursement from the out-of-state labs for the costs of inspections. By consistently failing to perform a sufficient number of biennial inspections—a core component of its oversight responsibilities—Laboratory Services has demonstrated a pattern of not ensuring that labs adhere to state requirements.
Laboratory Services Has Minimized Proficiency Testing Monitoring for Out-of-State Labs

According to Public Health’s assistant deputy director of the Office of the State Public Health Laboratory Director, Laboratory Services did not adopt new proficiency-testing policy and procedures (proficiency procedures) until March 2015—more than seven years after our 2008 audit. Further, it still has not addressed our recommendation in the 2008 audit that it promptly review the proficiency-testing results of out-of-state labs. Monitoring labs’ proficiency testing provides Laboratory Services with insight into labs’ performance between biennial on-site inspections.

Laboratory Services is responsible for monitoring proficiency testing for all state-licensed labs regardless of their geographic location. However, Laboratory Services’ proficiency procedures minimize its oversight of out-of-state labs. According to the examiner in charge of licensing out-of-state labs, there are currently around 450 such labs, which represent around 16 percent of the licensed labs required to participate in proficiency testing. However, Laboratory Services’ proficiency procedures do not adequately help it determine how many failures an out-of-state lab may have or when those failures may occur. Further, the proficiency procedures do not explain how Laboratory Services obtains out-of-state labs’ testing results: The approach Laboratory Services has developed produces reports of test results only for labs located within California. The proficiency procedures do detail that Laboratory Services will verify during each out-of-state lab’s license renewal that it is enrolled in proficiency testing and that it is enrolled to test its proficiency in all areas corresponding to the testing it performs. However, these functions are not the same as monitoring results from proficiency testing.

We did see evidence that Laboratory Services reviews the results of proficiency testing for out-of-state labs when it renews the labs’ licenses. Nevertheless, the proficiency procedures call for Laboratory Services to review the results every 30 days from in-state labs that reported results during that 30-day period. According to the clinical labs facility section chief of the Richmond office (facility section chief) who manages facility licensing including proficiency testing, the less frequent monitoring of out-of-state labs is not a problem because CMS agents in the states where these labs are located monitor them for CLIA purposes, so the risk that deficiencies will go undetected is low. Even so, Laboratory Services’ current responsibility is to know which out-of-state labs have failed proficiency testing so it can take appropriate action; its reliance on federal monitoring under CLIA highlights the redundancy of Laboratory Services’ oversight in this area.
Notwithstanding the limitations of its monitoring of proficiency testing for out-of-state labs, Laboratory Services has addressed certain aspects of our recommendations concerning its proficiency procedures for in-state labs. For example, the proficiency procedures it implemented in March 2015 specify how often it should review proficiency-testing results and what steps it should take to ensure that labs take timely corrective action, and they outline sanctions when labs do not correct problems. Additionally, we reviewed 10 testing failures for in-state labs that occurred in 2014 and found that Laboratory Services had identified them, contacted the labs, received responses within the time periods set in the proficiency procedures, and accepted the labs’ plans to resolve the deficiencies within reasonable periods of time.

Laboratory Services’ Complaint Procedures Still Need Enhancing

Laboratory Services has not successfully modified its complaint procedures in response to our prior audit. In 2008 we recommended that Laboratory Services address weaknesses in its complaint investigation practices and its related policy and procedures. Our recommendations were aimed at helping Laboratory Services track and prioritize complaints while also ensuring that substantiated allegations were corrected. Laboratory Services’ records show that it received an average of 177 complaints annually from 2008 through 2014. However, it still has not established time frames for completing complaint investigations, and some lower priority complaints may never be investigated. Finally, Laboratory Services has not defined in its procedures when its staff should revisit labs to verify that they have successfully corrected the most significant problems substantiated during complaint investigations.

Although seven years have passed since we recommended to Laboratory Services that it strengthen its complaint procedures, it has not adequately addressed all of our concerns. Specifically, we expected to find that Laboratory Services had established time frames to ensure that it completes complaint investigations promptly, but it has not done so. We reviewed Laboratory Services’ complaint logs from January 2014 through April 2015; these logs show that it received 218 complaints and that 13 were open as of May 2015. We reviewed five of these open complaints and found that Laboratory Services had not, in our view, promptly addressed two of them. Each of the two complaints alleged that the labs had not properly supervised unlicensed laboratory personnel, yet as of May 21, 2015, neither complaint had been closed or investigated. Laboratory Services received the first complaint in April 2014 and the second in October 2014; therefore, it had left complaints unresolved for 10 months to over a year.
When we inquired about these two complaints, the examiner tasked with performing the investigations stated that he had not done so because the investigations would involve several examiners observing the labs for extended periods of time. The examiner further explained that he did not ask his supervisor for approval to conduct extended observations or for staff support because he did not think the supervisor would approve his requests. The lack of documentation explaining why Laboratory Services did not perform these complaint investigations, along with the fact that management did not approve the examiner’s decision, highlights Laboratory Services’ informal complaint process in which examiners appear able to determine, without further management review, which complaints are worthy of investigation.

Laboratory Services’ complaint procedures also do not address our concerns from the 2008 audit regarding the receipt and prioritization of complaints. Specifically, the complaint procedures still allow any employee to accept complaints, which as noted in our 2008 audit, increases the risk that Laboratory Services will lose a complaint or overlook a matter of serious concern. The complaint procedures also state that the lowest priority complaints will be investigated at the next on-site inspection. Even though licensed labs do require biennial inspections, state law does not require registered labs to be routinely inspected. As a result, Laboratory Services is potentially leaving the lowest priority complaints it receives uninvestigated for up to two years for licensed labs and indefinitely for registered labs. Furthermore, because Laboratory Services does not inspect a high percentage of licensed labs, it may not investigate at all complaints it classifies as low priority.

Finally, Laboratory Services’ complaint procedures continue to lack detail regarding when its staff should ensure that labs take corrective action in response to completed investigations. In particular, the complaint procedures do not discuss when performing another on-site inspection is warranted to ensure that the offending lab has corrected significant deficiencies—those that place a patient’s health at risk. Although we reviewed five complaints that Laboratory Services had substantiated through its investigations and noted that it performed a follow-up on-site inspection in each case, the lack of clear guidance increases the risk that examiners may not verify a lab’s efforts to correct even the most egregious of cases. Even though Laboratory Services deserves credit for the follow-up inspections it performed, we continue to believe that it could enhance its policies by setting clearer expectations defining when its examiners should visit labs to verify that significant problems no longer exist.
Laboratory Services Has Failed to Strengthen Its Sanctions Activities

Laboratory Services has failed to respond to our 2008 recommendations that it guide staff in using its sanction authority, including ensuring that labs comply with sanctions by, for example, paying the imposed fines. We previously reported that Laboratory Services imposed 23 sanctions in the form of civil money penalties from 2002 through 2007; however, we identified only four facility-related sanctions that Laboratory Services imposed since our 2008 audit. Further, it did not collect any civil money penalties in fiscal years 2012–13 and 2013–14. With Laboratory Services having oversight of almost 2,800 licensed labs, we are skeptical that so few labs actually required sanctioning—such as civil money penalties or revocations of their licenses or registrations—in the last seven years. Overall, it appears that Laboratory Services’ sanctioning process suffers from inconsistent staffing and unreliable records regarding past sanctions activity, leading to Laboratory Services’ inability to determine whether it collected in full the civil monetary penalties it imposed. Further, we believe Laboratory Services needs to develop a more robust sanctioning process, such as one that involves multiple managers who monitor sanction activity and collections, in order to guard against the potential for fraud.

Laboratory Services’ former chief stated that since our 2008 audit Laboratory Services has not always had staff dedicated to sanctioning efforts, noting that the most recent manager responsible for sanctioning held that duty for just one year before retiring. We also found that Laboratory Services has not updated its sanction policies since 1998. Two sanction cases we reviewed illustrate Laboratory Services’ lack of adequate processes that would allow it to sanction labs effectively. In the first case, we found that Laboratory Services did not promptly sanction a lab for willful and unlawful conduct when it employed and used unlicensed lab personnel to conduct tests and analyses. Laboratory Services received a complaint about this particular lab in February 2014, and after performing an on-site inspection in March 2014, notified the lab in May 2014 that it had confirmed the allegation. Nevertheless, in May 2015—one year after it confirmed the wrongdoing—Laboratory Services was still drafting a letter indicating its intent to impose a monetary penalty on the lab. Laboratory Services further delayed issuing the sanction letter by a month when its former chief retired. It ultimately mailed the letter in June 2015 and notified the lab of its intent to impose a $14,150 civil money penalty, plus an additional $7,500 to recover the costs of its investigation. By not promptly sanctioning the lab in this case, Laboratory Services showed its ineffectiveness at ensuring that labs adhere to state requirements.

Laboratory Services lacks staff dedicated to sanctioning efforts and lacks up-to-date policies, which would allow it to sanction labs effectively.
In the second sanction case, Laboratory Services could not provide documentation that a sanctioned lab had paid its civil money penalty in full. Laboratory Services imposed a sanction on a high-profile lab for willful and unlawful conduct for not having a lab director responsible for operations, for employing an unlicensed person who performed complex tests, and for submitting false statements on lab-licensing documents. Following a legal dispute with Public Health about this sanction, the lab agreed to pay $40,000 in 40 monthly installments of $1,000 from February 1, 2009, through May 1, 2012. However, the accounting reports that Laboratory Services provided to us did not account for $5,000 of the $40,000 penalty, and the former chief stated that Laboratory Services searched its records and could not find documentation indicating whether the lab paid the remaining $5,000. She offered the explanation that any supporting document might have been destroyed per Public Health’s records retention policy. She also stated that Laboratory Services did not develop a final notice documenting that the lab paid the penalty in full. Although it is possible the lab paid the full amount, Laboratory Services’ inability to resolve our questions concerning the receipt of the $5,000 demonstrates that it needs to strengthen its record keeping.

Laboratory Services’ lack of assigned staff, outdated processes, and unreliable data leave its sanctioning process vulnerable to mistakes and susceptible to staff engaging in fraudulent acts. Although we found no evidence of fraud, we noted that only one staff person knew the status of outstanding sanctions and other staff were required to turn to this person for answers to our questions about the sanctions process. For example, Laboratory Services’ staff were unable to provide a sanction database that was purported to track civil money penalties since, according to an associate governmental program analyst, only the one staff member in question had access to the file.

Further, once Laboratory Services provided us with the sanction database, we found that the data it contained were inconsistent with Laboratory Services’ official accounting reports. For example, the database showed that one lab paid a $92,840 penalty in June 2011; however, the accounting reports showed the penalty payment was posted to Laboratory Services’ Clinical Laboratory Improvement Fund (fund) but then reversed, thus cancelling the entry. Laboratory Services was unable to provide documentation that demonstrated where and if the funds were ever deposited, and Public Health’s accounting unit did not respond to our requests for further documentation and clarification about the reversing entry. With sanction records that cannot be easily compared with official accounting records, and with access to sanction information generally limited to one staff person, Laboratory Services’ sanction...
process is at risk for fraud and abuse. Because it continues to face challenges in its sanction program seven years after our 2008 audit, it is clear Laboratory Services has not addressed the concerns we raised at that time.

**Management of the Laboratory Services Program Is Inadequate**

Laboratory Services has also failed to respond to the recommendations we made in our 2008 audit for it to better manage its resources; consequently, problems that existed more than a decade ago still plague it. We found that mismanagement caused Laboratory Services to collect improper fee amounts from labs, to waste opportunities to partner with accreditation organizations that could boost lab oversight, and to fail to address hiring and retention issues in the face of an aging workforce. Laboratory Services has also taken little or no action to address several other recommendations we made, including those to improve its disjointed information technology systems and to update its outdated regulations, as shown in Table 3.

**Table 3**

**Laboratory Field Services’ Implementation of the California State Auditor’s 2008 Recommendations Related to Its Management Responsibilities**

<table>
<thead>
<tr>
<th>MANAGEMENT RESPONSIBILITY</th>
<th>2008 FINDING</th>
<th>RECOMMENDATION</th>
<th>RECOMMENDATION’S CURRENT STATUS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fee Adjustment</td>
<td>Laboratory Services (Laboratory Services) incorrectly adjusted clinical laboratory (lab) fees for three of the five years analyzed.</td>
<td>Laboratory Services should adjust its fees in accordance with the State's annual budget act.</td>
<td>No action taken.</td>
</tr>
<tr>
<td>Accreditation Organizations</td>
<td>Laboratory Services had not approved any accreditation organizations to help it oversee labs and meet its mandate.</td>
<td>Laboratory Services should use accreditation organizations to help it perform inspections.</td>
<td>Not fully implemented.</td>
</tr>
<tr>
<td>Hiring and Succession Planning</td>
<td>Laboratory Services attributed its inability to meet its mandated responsibilities to a lack of resources. It specifically identified inadequate staffing as a concern.</td>
<td>The California Department of Public Health, in conjunction with Laboratory Services, should ensure that Laboratory Services has sufficient resources to meet all its oversight responsibilities.</td>
<td>Not fully implemented.</td>
</tr>
<tr>
<td>Information Technology</td>
<td>Laboratory Services’ information technology data systems did not adequately support its activities, such as tracking all aspects of its complaints and sanctions activities. The information technology systems lacked safeguards to ensure that data were accurate and could be used by management.</td>
<td>Laboratory Services should ensure that its information technology systems support its needs, and if it continues to use internally developed databases, it should develop and implement appropriate system controls.</td>
<td>No action taken.</td>
</tr>
<tr>
<td>Regulations</td>
<td>In three instances, Laboratory Services maintained state regulations that state law had superseded.</td>
<td>Laboratory Services should update, repeal, and revise its regulations as necessary.</td>
<td>No action taken.</td>
</tr>
</tbody>
</table>

Sources: California State Auditor’s (state auditor) Report 2007-040 and the state auditor’s analysis of Laboratory Services’ corrective action.
Laboratory Services Is Overcharging Licensed Labs

Laboratory Services has continued to fail to correctly adjust its license fees, which has resulted in it overcharging labs more than $1 million. State law requires Laboratory Services to adjust its license and registration fees by percentages specified in the State’s annual budget act; however, the Legislature created a sliding schedule for license fees in 2009, which raised the fees, and since then license fees have been excluded from the annual budget act adjustment. We identified three errors Laboratory Services made since our 2008 audit. Laboratory Services’ most egregious error occurred in January 2014 when it implemented an unauthorized license fee increase of more than 13 percent. We estimate this error resulted in labs collectively overpaying the State more than $1 million in fees. Further, since posting the increased license fee in January 2014, Laboratory Services has continued to charge labs these erroneous fee amounts. We determined that Laboratory Services did not realize that the percentage increase did not apply to license fees, as specifically stated in the annual budget act. Laboratory Services does not have well-defined processes in place to ensure that it analyzes annual budget act changes accurately. We recommended in our 2008 audit that Laboratory Services should work with Public Health’s budget section to ensure that it adjusts fees in accordance with the budget act, and we had concluded in September 2009 that it fully implemented this recommendation based on information it had provided to our office. However, the unauthorized increase we identified during our follow-up audit has caused us to conclude that Laboratory Services needs to take additional action, particularly in light of recent retirements of key staff. For example, our review of Laboratory Services’ procedures revealed only a high-level, one-page document that did not describe who was responsible for coordinating with Public Health’s budget section to ensure that Laboratory Services implemented the correct fee adjustment each year. Moreover, when we tried to obtain Laboratory Services’ perspective on the improper increase, the facility section chief informed us that she was not involved in the fee calculations and that the employee who prepared the calculations had retired from Laboratory Services. The facility section chief stated that Laboratory Services’ former chief oversaw the fee increase process; however, the former chief also had retired. Key staff retirements only reinforce the need for Laboratory Services to have a clearly defined and well-understood process for increasing fees, which would include identifying those staff responsible for coordinating with Public Health’s budget section and the steps for verifying that the proper authorization exists in the annual budget act to execute fee changes.
Further, Laboratory Services’ revenue from license and registration fees has far exceeded its oversight costs. For fiscal years 2008–09 through 2013–14, Laboratory Services collected a total of about $31.2 million in license and registration fees while spending only $18.6 million to monitor labs. As a result, Laboratory Services collected roughly $12.6 million in fee revenue that it did not need for the level of oversight it provided. Figure 5 on the following page shows that Laboratory Services’ revenue collection for its fund has consistently exceeded its expenditures. Laboratory Services’ excess license and registration revenue may be in part due to Public Health’s decision to sponsor legislation establishing higher fees. In sponsoring this legislation, it argued that Laboratory Services did not have the resources necessary to adequately enforce state law by conducting inspections and investigating complaints. In October 2009 the new law became effective, but as we describe elsewhere in this report, Laboratory Services continues not to meet its oversight mandates despite the increased fees as reflected in its growing fund balance.

Moreover, Laboratory Services’ problems with overcharging labs may extend to overcharging the lab personnel who also pay fees. To qualify to perform lab work, state law requires certain lab personnel be licensed by Laboratory Services, which charges fees for the licenses. In addition to paying a fee, an individual must meet educational and training requirements and pass examinations. As of June 30, 2014, the fund’s total ending balance—for lab license and registration and personnel licensing—exceeded $19.3 million. Although at least $12.6 million of that balance pertained to Laboratory Services’ overcharging of labs, the remaining amounts may be due to Laboratory Services’ excessive revenue from personnel fees.

According to Laboratory Services’ health program manager, who is responsible for managing Laboratory Services’ accounting unit, the disparity between Laboratory Services’ licensing revenue and expenditures relates to its unfilled examiner positions. However, this explanation differs from the explanation Laboratory Services provided during our 2008 audit. At that time we inquired about Laboratory Services’ fund balance with the former assistant deputy director of the Center for Health Care Quality (former deputy director) who oversaw Laboratory Services. The former deputy director asserted that Laboratory Services would use the excess money in its fund for one-time investments to help stabilize the program, such as replacing Laboratory Services’ information technology systems. However, seven years later, Laboratory Services continues to have vacant facility examiner positions and has yet to replace its information technology systems. Although we would expect Laboratory Services to maintain a prudent reserve, such as an amount equaling 5 percent For fiscal years 2008–09 through 2013–14, Laboratory Services collected $31.2 million in license and registration fees while spending only $18.6 million to monitor labs—collecting $12.6 million that it did not need for the level of oversight it provided.
of its annual expenditures or about $420,000, Laboratory Services’ reserve exceeded $18 million as of June 30, 2014. With such a high reserve, the labs that pay fees to Laboratory Services may reasonably question the State as to the fairness and appropriateness of those fees in relation to the actual expense incurred for oversight.

Figure 5
Laboratory Field Services’ Revenues, Expenditures, and Ending Fund Balances Related to Its Oversight of Clinical Laboratories
Fiscal Years 2008–09 Through 2013–14

However, resolving Laboratory Services’ excessive reserve may prove difficult. A gap exists in state law such that Laboratory Services lacks the authority to ensure that its lab fees are consistent with the costs of oversight. On one hand, according to statute, Laboratory Services may charge only the amounts needed to cover its costs; the law states that total fees collected shall not exceed the costs incurred for licensing, certification, inspection, or other activities relating to the regulation of labs. On the other hand, the
Legislature currently uses the budget act to annually prescribe license and registration fee adjustments. Consequently, our legal counsel has advised that Laboratory Services lacks the authority to reduce its fees when its revenues exceed its costs. Based on its high fund balance at the end of fiscal year 2013–14, we estimate that Laboratory Services could suspend lab fee collection for the next three years and still have money remaining in its fund. Nevertheless, it is unclear what steps Laboratory Services can take to better align revenue with costs, such as lowering its fees or temporarily suspending them. We believe that the fees a regulated community pays should align with actual costs. Further, when fee revenue greatly exceeds costs, it is prudent for the State to achieve equilibrium in the most expeditious and administratively simple way possible.

**Laboratory Services Has Failed to Partner Effectively With Accreditation Organizations**

Laboratory Services has wasted opportunities to work with accreditation organizations to help it fulfill its oversight responsibilities. Under state law, Laboratory Services can approve private nonprofit accreditation organizations to conduct oversight functions—including performing inspections and monitoring proficiency testing—in lieu of its direct oversight. However, Laboratory Services has not taken full advantage of this opportunity, despite an internal analysis showing the potential positive effects on its workload and on its ability to meet its mandates. In an internal memo to the former chief of Laboratory Services dated February 2015, the former section chief of the Los Angeles office noted that six different accreditation organizations have accredited labs operating in California. The former section chief concluded that approving the six accreditation organizations would have a major impact on Laboratory Services’ workload. For example, approving the six accreditation organizations would reduce the number of labs requiring inspection by 1,254 and create a staffing surplus at Laboratory Services. The internal memo also acknowledged that with a limited staff of three performing inspections, the Los Angeles office was completing only 39 percent of its required workload. However, our follow-up audit found that Laboratory Services has approved only one accreditation organization, and it lacks a documented agreement formalizing that organization’s responsibilities for monitoring labs’ compliance with California law.

Laboratory Services has also not developed a process to approve and oversee accreditation organizations. A 2009 law clarified Laboratory Services’ existing authority to use accreditation...
organizations to help it with its oversight mandate. The 2009 law outlined application requirements and gave Laboratory Services about 15 months to develop and implement a program. As seems reasonable with any new agency initiative, we expected Laboratory Services to develop a plan including policy and procedures for accepting, processing, and approving accreditation organizations’ applications. We also expected Laboratory Services to establish plans for monitoring accreditation organizations’ performance after approval, including processes to periodically reauthorize each accreditation organization and revoke authorization when necessary. Ideally, these plans would contain detailed safeguards—such as multiple application reviewers, set time frames, and performance measures—that would help ensure that the processes were consistent and effectively managed. The 2009 law specified that Laboratory Services could issue its plan to use accreditation organizations through an All Clinical Labs Letter, which would take effect 45 days following publication. Despite the potential for this alternative regulatory process to be more streamlined than the process that Public Health would typically have to follow to implement new regulations, Laboratory Services did not use the authority the 2009 law granted it.

With no formal application review process in place, we noted that the applications of other accreditation organizations have awaited Laboratory Services’ decisions far beyond the time frame established in statute. State law required Laboratory Services to begin accepting applications from accreditation organizations on January 1, 2011, and to make a determination within six months of their receipt. Although Laboratory Services started accepting applications on time, it has failed to make determinations on all but one of the applications it has received. Four accreditation organizations submitted applications, and Laboratory Services took 20 months to approve one. It was still reviewing two of the other applications as of April 2015, nearly four years later. According to Laboratory Services’ former chief, the fourth and final application was withdrawn. She stated that she and a retired annuitant accepted and reviewed applications until July 2014, when the retired annuitant left Laboratory Services. At that time, the former chief assumed sole responsibility for reviewing and approving the accreditation organizations’ applications until her own retirement in May 2015. After the former chief retired, the assistant deputy director stated that she transferred the review of the remaining applications to two CLIA section staff, including the section chief, with the goal of responding to the accreditation organizations by the end of July 2015. However, when we followed up with the assistant deputy director at the end of July 2015, she stated that Laboratory Services would not meet the goal for responding to the
accreditation organizations due to the sheer volume of information in the application packets and that it did not have a date for potential approval.

For the one approved accreditation organization, Laboratory Services has not established or documented clear expectations for how it will monitor the organization's performance and transfer its oversight responsibilities. Although it granted the accreditation organization approval in August 2013, Laboratory Services has not entered into an agreement with it specifying the organization's role and responsibilities or establishing how Laboratory Services, through oversight, will verify that it is performing acceptably. When we inquired about the status of Laboratory Services drafting and signing agreements with approved accreditation organizations, we received conflicting viewpoints. The assistant deputy director recognized that relying on accreditation organizations for lab oversight involved some risk, and she stated that she would be more comfortable if a memorandum of understanding existed between the approved accreditation organization and Laboratory Services. In contrast, the former chief stated she believed that the existing statute outlines the responsibilities of the accreditation organization, thereby implying that additional formal agreements were unnecessary.

Despite its internal analysis outlining the benefits of using accreditation organizations for oversight, Laboratory Services has not capitalized on this opportunity. Specifically, it has no guidelines to review and approve accreditation organizations’ applications, no documented plans to monitor their performance, and no formal agreement to ensure that responsibilities are clearly articulated to facilitate accountability. With Laboratory Services' inspection responsibilities largely unmet, we find it surprising that it has not prioritized using accreditation organizations.

Laboratory Services Faces Significant Staffing Challenges and Has Failed to Plan for Retirements Through Succession Planning

Over the past seven years, Laboratory Services has not resolved the issues that it claimed have kept it from having sufficient staffing to meet its mandate. For example, during our 2008 audit, Laboratory Services explained that it did not plan to conduct regular inspections of labs every two years unless it received additional resources, noting at the time that it had only three examiners focused on investigating complaints and inspecting labs for initial licensure. In January 2014 Laboratory Services’ management drafted a recruitment and retention proposal that aimed to increase employee salaries and thus make the examiner position more attractive to both future and current employees. However, that
Laboratory Services claims that raising salaries will improve its ability to hire and retain staff. However, it has yet to convince its human resources branch that it has a compelling argument for requiring higher salaries.

Laboratory Services claims that raising salaries will improve its ability to hire and retain staff. In responding to the recommendations from our 2008 audit, Laboratory Services’ management repeatedly stated that salaries were a barrier to staff recruitment and retention. In its January 2014 proposal, Laboratory Services’ management wrote that the salaries for its examiner staff had lagged behind the private sector for years and if Public Health did not approve the proposal, the salary imbalance would cause Laboratory Services to continue to lose qualified examiner staff. The proposal further warned that the lax oversight that would inevitably result from the overextension of its shrinking examiner staff would compromise Laboratory Services’ ability to assure high-quality laboratory testing and health care for the people of California.

However, Laboratory Services has yet to convince the human resources branch within in its own department that it has a compelling argument for requiring higher salaries. The chief of Public Health’s human resources branch (human resources chief) and her staff reviewed the proposal and responded in January 2014, concluding that Laboratory Services’ proposal lacked compelling evidence for the requested salary increase. Although not disputing the need for more examiners, the human resources chief concluded that Laboratory Services already had sufficient resources to fill its current vacancies, noting for example that it had 43 candidates on its hiring lists for six open examiner positions. She stated that Laboratory Services needed to demonstrate through evidence that it either sent the individuals on these hiring lists contact letters and they waived their interest in employment or that it interviewed them and found them to be unsuitable for employment. According to the human resources chief, Laboratory Services could not defend its contention that it had a recruitment problem without such documentation. She was equally skeptical that Laboratory Services had a retention problem, noting that since July 1, 2007, only one examiner had resigned and one failed probation, with the remaining separations resulting from retirements. Summing up her evaluation of the proposal, she
stated that Laboratory Services needed to do more than just make a request and issue statements; it had to defend its position with more data.

Before Laboratory Services had even made an internal proposal to increase salaries for its examiners in 2014, the Legislature authorized it to hire 16 new examiners for lab facility oversight pursuant to the 2010 Budget Act (Chapter 712, Statutes of 2010). However, according to the former chief, it was unable to fill the new positions the Legislature approved because the governor implemented a hiring freeze in February 2011, four months after the 2010 Budget Act was passed and Laboratory Services gained the authorization to hire. Although it appears to us that Laboratory Services may have been eligible for an exemption from the hiring freeze based on the criteria set forth by the governor’s office, the former chief only provided documentation supporting one exemption request for one examiner position focused on lab facilities. Given Laboratory Services’ long-standing claims that it needed additional staff to meet statutory requirements for lab oversight, and given that funding for this oversight comes from the fees that labs pay, we believe Laboratory Services could have made a strong case for an exemption from the governor’s hiring freeze, thus increasing the number of examiners dedicated to lab oversight.

Today, Laboratory Services has a significant number of examiners approaching retirement, yet it has not developed a succession plan to confront this problem. Based on an analysis Laboratory Services performed, the average age of a Laboratory Services examiner in 2013 was roughly 61, with management-level examiners having an average age of 65. We expected to find Laboratory Services had adopted a succession plan that would include identifying staff competency gaps, developing strategies to address those gaps, and identifying and developing the potential of current employees to fill key leadership positions. However, Laboratory Services’ management has not developed or implemented a succession plan. The assistant deputy director stated that Laboratory Services has handled succession planning by bringing back retired annuitants and that historically Laboratory Services has not developed staff through training to prepare them to move into senior positions. She also acknowledged that succession planning is important because a significant number of staff are eligible for retirement. She said she is currently drafting a reorganization plan that accounts for staffing and succession difficulties; however, the timing of that plan’s approval and implementation is uncertain. Although the assistant deputy director asserted she would like to start implementing her plan in October 2015, she noted that the plan hinges on approval from Public Health’s human resources branch and is dependent on previously lost positions being reestablished.
Laboratory Services has not updated or substantially improved its information technology systems to adequately support its activities.

Laboratory Services has not updated or substantially improved its information technology systems to adequately support its activities. Table 3 on page 25 summarizes the information systems issues we identified in 2008. When an information technology system contains illogical, incomplete, or incorrect data, its usefulness as a tool to aid management’s decision making is limited. Laboratory Services’ continued reliance on information technology systems containing flawed data shows that it has not taken steps to address the recommendations from our 2008 audit.

The information technology systems relevant to Laboratory Services include the Health Applications Licensing system (HAL), which is a legacy system that provides licensing information for labs. Although seven years have passed since our 2008 audit, Laboratory Services has not improved HAL in response to our audit recommendations. Public Health’s information technology services division provided us with a log of changes it made to HAL since 2008 in response to Laboratory Services’ requests, which included changes such as exporting a list of lab directors to a downloadable file rather than to a printer and making modifications to individual lab records. However, none of the changes addressed our recommendations. For example, in our 2008 audit, we found that the complaint field was limited to a yes or no indicator and that HAL’s lack of additional fields for information such as the nature or status of the complaint limited the system’s usefulness as a management tool. Nevertheless, the change logs do not reflect that Laboratory Services requested additional fields for recording complaint information in HAL.

Laboratory Services also uses four Microsoft Access databases to track complaints and sanctions. We found that the problematic conditions with these databases that we identified in our 2008 audit were generally unchanged, confirming that Laboratory Services has not responded to our recommendations. Specifically, we were unable to verify that many of the complaints appearing in the complaint logs were listed in the complaints database. Also, the sanction database did not include an ongoing sanction concerning a lab that employed unlicensed personnel to perform highly complex testing. In the most striking example, in 2013 Laboratory Services received more than 100 complaints according to its complaint logs, but none of these complaints were recorded in its complaint database for that year. Because it does not consistently track complaints in its database, Laboratory Services is forced to research how many complaints are open at any given time. Further, the paper complaint logs we reviewed did not consistently include information about the nature of the specific complaints received,
thus making more difficult Laboratory Services’ task of prioritizing complaint investigations or evaluating the frequency and types of allegations against specific labs.

Laboratory Services does not have specific plans to upgrade the information technology systems relevant to its work, and the assistant deputy director could not provide a time frame for when Public Health might consider such plans. She explained that although Public Health wants to improve Laboratory Services’ information technology systems, it has not yet begun work on the necessary feasibility studies. She said that Public Health will not begin these studies until it completes another information technology system project, and she could not provide a date for that.

Laboratory Services has also not updated lab regulations since our prior audit, although we found that it has identified lab regulations that it needs to change. In our last report, we identified specific regulations that were not consistent with state law. For example, state regulations define unsuccessful participation in proficiency testing as three consecutive failures, while state law, as amended to adopt federal regulations, defines unsuccessful participation as two consecutive failures or two failures out of three consecutive tests. We expected to find that Laboratory Services had taken action to repeal outdated state regulations, thereby averting misunderstandings both within Laboratory Services and between it and the regulated community. Nonetheless, Laboratory Services has not taken action to change the regulations. An attorney with Public Health provided us a log she asserted Public Health uses to track its regulatory packages; the log reflects that Public Health plans to submit five regulations packages related to labs to the California Secretary of State from March 2016 through December 2019.

The State’s Oversight of Clinical Labs Largely Duplicates Efforts at the Federal Level, Raising Questions as to Whether a Separate State Approach Is Needed

With Laboratory Services’ history of failing to perform its oversight responsibilities, the State has, in effect, relied on CLIA to ensure that labs perform accurate testing. Even if Laboratory Services were fulfilling its mandates, the core requirements found in state law concerning the licensing and oversight of labs duplicate those found in CLIA. The duplication does not appear to provide any added benefit to California’s consumers, in part because Laboratory Services has historically been unable to manage its workload, as discussed earlier in this report, and in part because the requirements set out in CLIA and monitored by CMS represent a reasonable alternative to Laboratory Services’ failed oversight.
We believe the Legislature should consider eliminating the requirement that the State license labs and instead rely on CLIA and the licensing and oversight structure CMS manages, as many other states do. We believe that eliminating the duplicate state requirements would have a negligible effect on the CLIA section’s workload because the bulk of its responsibility is inspecting and monitoring proficiency testing for CLIA-certificated labs, and the number of these labs would not change.

As previously mentioned, Laboratory Services also currently issues and monitors licenses for personnel who work in labs to ensure they meet state requirements. We did not review Laboratory Services’ effectiveness at administering the state licensing requirements for lab personnel; therefore, we believe Laboratory Services should maintain this responsibility at this time.

The State Law That Specifies Lab Requirements Largely Duplicates CLIA

The state law that mandates Laboratory Services’ oversight of labs largely duplicates CLIA’s requirements. Given the significant similarities, along with Laboratory Services’ difficulty in completing its oversight responsibilities, we question the State’s need to maintain lab requirements separate from CLIA’s. The core state and federal requirements for licensing and oversight of labs are summarized in Table 4 and, as the table shows, the requirements in state law and CLIA are identical. For example, both state law and CLIA would require labs that conduct complex tests on specimens originating in California, to be authorized to perform tests and to pay fees for oversight. Both state law and CLIA would also require that labs receive ongoing, periodic oversight composed of biennial inspections and monitoring of their proficiency-testing results. Finally, both state law and CLIA provide for complaint investigations and sanctions of labs as needed to ensure that they correct any deficiencies.

Lab requirements and the oversight embodied in state law reach beyond California’s borders when out-of-state labs test samples originating from within the State. Thus, labs in other states and countries must obtain licenses or registrations from Laboratory Services if they test specimens from California. CLIA applies to labs in a similarly broad manner; all labs, including those in other countries, that test specimens collected in the United States and its territories are subject to CLIA. Therefore, as Figure 6 on page 38 shows, all labs analyzing specimens originating in California are subject to state law and to CLIA. For example, a lab operating in Michigan that performs complex tests on samples originating in California would be subject to Laboratory Services’ biennial
inspections, proficiency testing, and oversight, and it would also be bound to the requirements found in CLIA as monitored and enforced by CMS and its agents.

Table 4
A Comparison of the Core Requirements in State Law and the Clinical Laboratory Improvement Amendments of 1988

<table>
<thead>
<tr>
<th>CORE REQUIREMENT</th>
<th>STATE LAW</th>
<th>CLINICAL LABORATORY IMPROVEMENT AMENDMENTS OF 1988 (CLIA)</th>
</tr>
</thead>
<tbody>
<tr>
<td>All clinical laboratories (labs) must be authorized to analyze specimens.*</td>
<td>✔</td>
<td>✔</td>
</tr>
<tr>
<td>All labs must pay a fee for initial authorization and then periodically to renew.†</td>
<td>✔</td>
<td>✔</td>
</tr>
<tr>
<td>Labs must be inspected biennially.‡</td>
<td>✔</td>
<td>✔</td>
</tr>
<tr>
<td>Labs must enroll and successfully participate in proficiency testing.‡</td>
<td>✔</td>
<td>✔</td>
</tr>
<tr>
<td>Complaints against labs are investigated, which may include on-site inspections.‡</td>
<td>✔</td>
<td>✔</td>
</tr>
<tr>
<td>Labs may be sanctioned for failing to meet requirements.‡</td>
<td>✔</td>
<td>✔</td>
</tr>
</tbody>
</table>

Sources: California Business and Professions Code; Title 42, Code of Federal Regulations, Section 493; and auditor analysis of state and federal law.

* State law requires labs to either register or obtain licenses depending on the types of tests they perform. CLIA requires labs to apply for one of several certificates depending on the types of tests they perform.
† State law requires labs to renew their licenses or registrations annually. CLIA requires labs to apply for new certificates biennially.
‡ Labs performing moderately complex to highly complex tests are subject to biennial inspections and proficiency testing. Labs performing simple tests are not subject to these two requirements. However, all labs are subject to complaint investigations and sanctions.

Under CLIA, states are allowed to develop their own licensing programs and requirements that are more stringent than federal standards. Once it adopts such requirements, a state can request that CMS exempt it from CLIA’s requirements and thus retain full oversight over the labs within its jurisdiction. Currently, only two states—Washington and New York—are exempt from some or all of CLIA’s requirements. In the mid-1990s, California considered applying for its own CLIA exemption. The Legislature passed Senate Bill 113 (Chapter 510, Statutes of 1995) to make several changes to state law in an attempt to incorporate CLIA’s standards while enacting more stringent standards for lab personnel, thus placing the State in a position to seek CLIA exemption. According to a report Laboratory Services prepared, the State earned CLIA exemption in 1999 but subsequently declined it because of concerns with paying an overhead fee—$2.4 million per year—to the federal Department of Health and Human Services. As a result, the State has been operating under a largely duplicate set of state and federal standards ever since.
Figure 6
Redundancy Between State and Federal Oversight Requirements for Clinical Laboratory Facilities

Source: California State Auditor’s analysis of state law and federal regulations.

* Labs performing moderate to high complexity tests—tests with a higher chance of risk or error such as hepatitis testing—are subject to state oversight inspections no less than once every two years. In contrast, state law exempts registered labs—those performing simpler tests with less chance of error such as prepackaged manufactured tests—from routine inspections, but the State is authorized to inspect them at any time it sees fit.

† Proficiency testing is the external evaluation of the accuracy of a lab’s test results. Only labs performing moderate to high complexity tests are subject to proficiency testing under state law and federal regulation.

‡ For state purposes, out-of-state labs are those labs located outside of California that test specimens originating from within California. These labs are subject to both California’s lab facility requirements and federal requirements.

CLIA Is a Reasonable Alternative to Laboratory Services’ Ineffective Oversight of Labs

Given Laboratory Services’ performance and management problems and the duplicate oversight structures that exist under state law and CLIA, it does not appear to us that California’s consumers receive meaningful protections from Laboratory Services’ oversight of labs. If the Legislature desires to eliminate the inefficient duplication, it could repeal state law requiring that labs obtain state licenses or registrations while leaving in place the State’s more stringent requirements governing lab personnel. In fact, most states do not have their own lab licensing programs and rely instead on the oversight structure that CMS administers. Based on an analysis an accreditation organization prepared, about 30 states—or just over 60 percent—did not have state lab-licensing
and oversight programs as of June 2014. In particular, some states with large populations—such as Texas and Michigan—do not issue state licenses to labs and instead work with CMS to enforce CLIA requirements. As a result, these states rely solely on CLIA oversight and the related monitoring and enforcement that CMS and its state agents provide.

CMS’s oversight process focuses on clinical labs being inspected and monitored by its state agents or accreditation organizations, while it monitors the oversight work these groups provide. As described in the Introduction, Laboratory Services acts as CMS’s state agent in California and has established a specific unit—the CLIA section—to provide oversight. According to CMS data, as of July 2015 the CLIA section oversaw 1,550 CLIA-certificated labs that performed moderately to highly complex tests. CMS uses a variety of means to help ensure that the CLIA section fulfills its oversight role. For example, CMS annually contracts with Laboratory Services and provides funding for CLIA section staff to participate in mandatory training. CMS also assesses the CLIA section’s performance through monitoring surveys that evaluate examiners’ performances while inspecting particular labs. The purpose of these monitoring efforts is to alert CMS if Laboratory Services’ staff require further training or other feedback as they monitor labs on behalf of CMS.

CMS also annually assesses the CLIA section through comprehensive performance evaluations. The CLIA section’s last two performance evaluations documented that it had met CMS’s expectations and developed and adhered to corrective action plans as necessary. Specifically, for the most recent evaluation in 2015, CMS commended the CLIA section for its fine performance because it exceeded CMS’s expectations for all criteria reviewed; consequently, no corrective action plan was necessary. The evaluation documented the CLIA section’s historical performance for certain oversight responsibilities and noted that since 2007 the CLIA section has earned perfect scores related to its proficiency testing process and complaints process. The CLIA section’s May 2014 performance evaluation showed that it was responsive to CMS, which had identified two labs during the prior year’s evaluation that it had not inspected in a timely manner. In response, the CLIA section developed a written corrective action plan detailing how it would ensure that it identified for inspection labs with expiring certificates, and CMS did not identify this issue in the CLIA section’s following year’s performance evaluation.

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2 According to CMS, 1,550 clinical labs were operating in California with a CLIA Certificate of Compliance in July 2015. These labs were not accredited by independent organizations. An additional 1,236 accredited labs operated in California with a CLIA Certificate of Accreditation. The CLIA section performs limited monitoring of the accredited labs at the direction of CMS.
CMS also takes steps to ensure that accreditation organizations maintain strict standards and fulfill their federal oversight role. In California, accreditation organizations oversaw 1,236 labs as of July 2015, according to CMS’s data. To become an authorized accreditation organization, an entity must provide CMS with a detailed comparison of its requirements and CLIA’s requirements and must describe its inspection process, its process for monitoring proficiency testing, and its process for responding to complaints. At least every six years, an accreditation organization must reapply to CMS to maintain its status as an authorized accreditation organization. In between application reviews, CMS requires its state agents to oversee the accreditation organizations by annually evaluating a subset of their lab inspection results; CMS may also conduct on-site inspections. For example, a state agent on CMS’s behalf will reinspect a lab following an accreditation organization’s inspection in order to compare inspection results; in California, the CLIA section performs these reviews on CMS’s behalf. If the state agent’s review shows substantial disparities from the expected results, CMS may terminate the accreditation organization’s ability to act as its agent.

As discussed in the Introduction, the director and assistant director from the Office of the State Public Health Laboratory Director oversee Laboratory Services. When we asked for their perspective on phasing out certain Laboratory Services mandates, they acknowledged that they have considered changes to Laboratory Services’ responsibilities, such as seeking to end the State’s licensing of labs in favor of following a CLIA-only model. However, the assistant director stated that they had concerns about the effect on Laboratory Services’ personnel licensing mandates if the Legislature eliminated its lab-licensing mandates. According to the director, the ability to inspect labs has given Laboratory Services a venue to enforce its personnel licensing requirements, but he acknowledged that Laboratory Services may simply need clear authority to enter facilities for personnel licensing enforcement. Ensuring that professionals have adequate training and experience to qualify them for laboratory positions is a valuable safeguard. Our proposal to eliminate lab-licensing and oversight requirements does not extend to personnel licensing requirements. We did not review Laboratory Services’ effectiveness at administering personnel licensing; therefore, at this time, it is our intention that those requirements be maintained.

Although the State has historically maintained lab requirements separate from CLIA, we believe that eliminating the duplicate requirements will have a negligible effect on the CLIA section’s workload. Most of the CLIA section’s current responsibilities will remain the same if the State discontinues its lab requirements. The CLIA section would continue to inspect nonaccredited labs,
its current responsibility. In contrast, Laboratory Services would cease to monitor accredited labs for compliance with state law; however, these accredited labs would continue to be monitored by their accreditation organizations and would still be subject to review by CMS or its agents in response to substantial allegations of noncompliance with federal requirements. The CLIA section’s focus would be CLIA-certificated labs; accredited labs would be overseen by the accreditation organization. The CLIA section might become responsible for additional complaints and sanctions if the State discontinues its lab requirements because complainants would no longer be able to file complaints with Laboratory Services. Thus, complainants who in the past might have filed complaints with Laboratory Services would need to file them with either the CLIA section or the accreditation organizations, depending on the labs’ authorization. For example, based on Laboratory Services’ pattern of assigning complaints for investigation, we determined that 69 of the 218 complaints Laboratory Services received between January 2014 and April 2015 might have required the CLIA section’s attention.

When we spoke with the CLIA section chief (CLIA chief), she indicated that California could become a CLIA-only state with minimal to no effect on the CLIA section’s workload. The CLIA chief noted that in the absence of a state-licensing requirement for labs, some complaints could be filed with her section. The CLIA chief clarified that under CLIA rules, CMS would direct complaints alleged against an accredited lab to the accreditation organization, but at times it might involve the CLIA section. The CLIA chief also shared that complaints that rise to the level of sanctions would require the CLIA section to prepare sanction proposals for CMS’s consideration. However, the CLIA chief confirmed that the CLIA section’s workload associated with inspections and proficiency-test monitoring would not change.

Employing accreditation organizations as oversight partners is a strategy CMS uses at the federal level, and the Legislature has long recognized the valuable role accreditation organizations can play in state oversight. With the passage of Senate Bill 744 (SB 744) (Chapter 201, Statutes of 2009), the Legislature provided for an application process whereby accreditation organizations, if approved by Laboratory Services, would have their accredited labs deemed as meeting the State’s requirements. Under this framework, the Legislature required that accreditation organizations inspect their member labs and monitor their proficiency testing in lieu of Laboratory Services performing these oversight duties. Although the Legislature maintained Laboratory Services’ authority to investigate complaints against accredited labs—which under federal rules is a responsibility left to the accreditation organizations—the oversight model the Legislature envisioned in SB 744 places several
of the core oversight responsibilities over clinical labs, such as performing inspections and monitoring the results of proficiency testing, in the hands of accreditation organizations.

Even though state and CLIA lab requirements are mostly the same, state law includes a few requirements that are not found in CLIA. One such requirement is that the lab owners must be disclosed on the State’s lab application; if the Legislature eliminates the requirement that labs obtain state licensure or registration, then the State would no longer have that information. According to the facility section chief of the Richmond branch, Laboratory Services uses owner information, for example, to ensure that owners who have operated labs whose licenses have been revoked or that have received sanctions cannot own or operate labs for two years. She also stated that CMS sometimes requests lab ownership information because the federal government does not track this information. However, we found that CMS and its agents obtain ownership information in different ways. For example, federal law requires that a lab owner or an authorized representative sign the CLIA certificate application and report any ownership change within 30 days after it occurs. Moreover, the CLIA certificate application requires the applicant to report his or her federal tax identification number, which could be used to obtain information about lab ownership. Finally, federal law also states that no person who has owned or operated a lab that has had its certificate revoked may, within two years, own or operate a lab operating under CLIA; to help enforce this law, CMS publishes an annual report on labs and persons who have violated CLIA requirements.

Another unique state requirement is that labs must conspicuously post their state licenses in their lab facilities; however, if the State no longer issues licenses, this requirement becomes moot. The facility section chief said she believes posting the license is for the public’s benefit because it promotes transparency and includes information that allows customers to know the lab is legitimate. However, we believe the Legislature could, for example, easily require that labs post their CMS-issued certificates to continue to promote public transparency. As a result, California’s unique state requirements should not pose a barrier to transitioning to a CLIA-only model.
Recommendations

Legislature

To eliminate the State's redundant and ineffective oversight of labs and to ensure labs do not pay unnecessary or duplicative fees, the Legislature should do the following:

- Repeal existing state law requiring that labs be licensed or registered by Laboratory Services and that Laboratory Services perform oversight of these labs. Instead, the State should rely on the oversight CMS provides.

- Repeal existing state law requiring labs to pay fees for state-issued licenses or registrations.

If the Legislature decides to continue requiring that clinical labs be licensed or registered through the State, it should amend state law establishing how Laboratory Services annually adjusts its fee amounts to ensure the revenue it collects does not exceed the cost of its oversight. Such an amendment might authorize Public Health to temporarily suspend or reduce fees when the Clinical Laboratory Improvement Fund's ending balance exceeds a prudent reserve amount that the Legislature establishes.

Regardless of whether it decides to repeal existing law, the Legislature should direct Laboratory Services to advise it on how best to address the millions of dollars in the Clinical Laboratory Improvement Fund in excess of a prudent reserve.

Laboratory Services

While the Legislature considers eliminating the requirement that labs obtain state-issued licenses or registrations and receive oversight from Laboratory Services, Laboratory Services should begin taking action to address its deficiencies by developing a corrective action plan by December 31, 2015. The corrective action plan should address its plans for implementing the recommendations from our 2008 audit and from this follow-up audit. For each item in its corrective action plan, Laboratory Services should identify the individuals responsible for ensuring it takes the corrective action, the resources it needs to carry out the corrective action, and the time frame in which it expects to successfully complete the corrective action.
To ensure it can provide effective oversight of labs as state law requires, Laboratory Services should do the following:

- Every two years, inspect all in-state and out-of-state labs it has licensed.

- Develop and implement proficiency testing policy and procedures for ensuring that it can promptly identify out-of-state labs that fail proficiency testing.

- Improve its complaints policy and procedures to ensure that it either investigates allegations promptly or clearly documents its management’s rationale for not investigating. It should also establish clear expectations for when staff must visit a lab to verify successful corrective action.

- Dedicate multiple staff to sanctioning efforts and update its sanctioning policy and procedures, including identifying steps to ensure that labs adhere to sanctions and that it collect civil money penalties. In addition, it should develop a single sanctions tracking system that multiple managers can monitor and that will allow it to periodically reconcile the monetary penalties it receives with Public Health’s accounting records.

- Work with Public Health’s budget section and other appropriate parties in developing a process to assess the budget act annually and to adjust its fees accordingly. The process should include its management’s review and approval of fee adjustments before it posts those fees publicly.

- Maximize the opportunity to partner with accreditation organizations by developing an accreditation organization program and issuing an All Clinical Laboratories Letter detailing the program’s components. In addition, it should consult with legal counsel and draft an agreement outlining the role and the responsibilities that Laboratory Services and the accreditation organizations will assume.

- Address staffing issues by preparing and resubmitting to Public Health a recruitment and retention proposal, developing a succession plan, and taking necessary steps to implement its planned reorganization.

- Ensure that its information technology data systems have necessary safeguards, contain accurate and complete data, and support its program needs.

- Update and develop its regulations as necessary to ensure consistency with existing state law.
We conducted this audit under the authority vested in the California State Auditor by Section 8543 et seq. of the California Government Code and according to generally accepted government auditing standards. Those standards require that we plan and perform the audit to obtain sufficient, appropriate evidence to provide a reasonable basis for our findings and conclusions based on our audit objectives specified in the Scope and Methodology section of the report. We believe that the evidence obtained provides a reasonable basis for our findings and conclusions based on our audit objectives.

Respectfully submitted,

ELAINE M. HOWLE, CPA
State Auditor

Date: September 10, 2015

Staff: Grant Parks, Audit Principal
      Sharon L. Fuller, CPA
      Kathryn Cardenas, MPPA
      Taylor William Kayatta, JD, MBA

Legal Counsel:  Richard B. Weisberg, Sr. Staff Counsel

For questions regarding the contents of this report, please contact
Margarita Fernández, Chief of Public Affairs, at 916.445.0255.
Blank page inserted for reproduction purposes only.
August 24, 2015

Elaine M. Howle  
State Auditor  
California State Auditor  
621 Capitol Mall, Suite 1200  
Sacramento, CA 95814

Dear Ms. Howle:

Enclosed is the California Department of Public Health’s (CDPH) response to the California State Auditor draft report titled “Follow-up: California Department of Public Health: Laboratory Field Services Is Unable to Oversee Clinical Laboratories Effectively, but a Feasible Alternative Exists” Report 2015-507, September 2015.

Thank you for the opportunity to respond. If you have questions please contact Monica Vazquez, Chief, Office of Compliance at (916) 440-7387.

Sincerely,

Karen L. Smith, MD, MPH  
Director & State Health Officer
California Department of Public Health Response
to Draft Report: “Follow-up: California Department of Public Health: Laboratory
Field Services Is Unable to Oversee Clinical Laboratories Effectively, but a
Feasible Alternative Exists”
Report 2015-507 September 2015

Recommendation 1
Inspecting licensed labs within and outside of California that test samples that
originate within the state every two years.

Response 1
The California Department of Public Health (CDPH) agrees with this recommendation
and is in the process of implementing. While our compliance rate for reviewing in-state
non-accredited laboratories has been high, Laboratory Field Services (LFS)
acknowledges that our overall compliance rate must improve. To that end, LFS is
analyzing its business processes, policies and procedures to ensure efficiencies and
reduce redundancies. LFS will focus specific personnel on clear tasks and streamlined
processes will increase the number of inspections performed. In addition to improving
existing processes, LFS has approved one accrediting organization (AO) that can
perform inspections on behalf of the State, and is currently reviewing two additional AO
applications. Approval of AOs will significantly decrease onsite workload and will
facilitate LFS meeting its mandated workload.

LFS, in partnership with CDPH’s Human Resources Branch (HRB), will increase its
recruitment and retention efforts to hire and maintain staff who can assist with this effort.

LFS will have a corrective action plan developed by December 31, 2015 to address this
recommendation.

Recommendation 2
Developing and implementing proficiency testing policy and procedures for
ensuring that it can promptly identify out-of-state laboratories that fail proficiency
testing.

Response 2
CDPH agrees with this recommendation. As a result of the audit released in 2008, LFS
updated its proficiency testing policies and procedures and implemented this revision in
March 2015. As a result, in-state laboratory proficiency testing has improved. However,
LFS will continue to update the out-of-state laboratory proficiency testing policies and
procedures. These updates will address the California State Auditor’s (CSA) concerns
and improve the rate at which LFS monitors and responds to proficiency testing results
of out-of-state laboratories. A corrective action plan will be developed by December 31,
2015.
California Department of Public Health Response
to Draft Report: “Follow-up: California Department of Public Health: Laboratory Field Services Is Unable to Oversee Clinical Laboratories Effectively, but a Feasible Alternative Exists”
Report 2015-507 September 2015

Recommendation 3
Improving its complaints policy and procedures to ensure that allegations are either investigated promptly or that management’s rationale for not investigating is clearly documented, and to establish clear expectations for when staff must visit a laboratory to verify successful corrective action.

Response 3
CDPH agrees with this recommendation. As a result of the audit released in 2008, LFS updated its complaints policies and procedures and implemented this revision in March 2015. Further, LFS established a complaints investigator position, and this person started employment in August 2015. LFS will improve its monitoring system to track complaints and perform audits to ensure complaints are timely completed and properly documented. When possible and applicable, LFS coordinates complaints with field surveys to efficiently utilize staff, and will enhance the complaint policies and procedures to address complaints for registered facilities. We are currently reviewing the complaints policies and procedures to address CSA’s concerns and will have a corrective action plan in place by December 31, 2015.

Recommendation 4
Dedicating multiple staff to sanctioning efforts and updating its sanctioning policy and procedures, including steps to ensure that sanctions are adhered to and civil money penalties are collected. In addition, it should develop a single sanctions tracking system that multiple managers can monitor and with which monetary penalties received can be periodically reconciled with Public Health’s accounting record.

Response 4
CDPH agrees with this recommendation and is in the process of implementing. As a result of the audit released in 2008, LFS is in the process of updating its enforcement and civil money penalties policies and procedures. The updates will allow us to determine the number of staff needed to oversee our sanctioning efforts and update the tracking system as required. LFS will establish a system to track sanctions, monitor issuance and collection of penalties, reconcile penalty payments with our accounting records, and allow staff to perform compliance audits necessary. LFS will have a corrective action plan in place by December 31, 2015 that will address all issues in this recommendation.

Recommendation 5
Working with Public Health’s budget section and other appropriate parties in developing a process to assess the budget act annually and to adjust fees accordingly, including management’s review and approval of fee adjustments before those fees are posted publically.

Response 5
CDPH agrees with this recommendation. LFS will work closely with our Budget Division to ensure annual fee adjustments are accurately calculated. In partnership with CDPH’s Administration Division and Office of Legal Services, LFS will also develop policies and procedures for calculating annual fees, including fee adjustments, and ensuring these fees are posted appropriately. These policies and procedures will help ensure consistent application of fee increases in the event of staff retirement or separation. A corrective action plan will be developed by December 31, 2015.

Recommendation 6
Maximizing the opportunity to partner with accreditation organizations by developing an accreditation organization program and issuing an All Clinical Laboratories Letter detailing the program’s components. In addition, consulting with legal counsel and drafting an agreement outlining the role and the responsibilities Laboratory Services and an accreditation organization will assume.

Response 6
CDPH agrees with this recommendation and is in the process of implementing. CDPH has posted on its LFS website criteria for submission for approval by CDPH of AOs. As a result of that posting, three AOs have applied for deeming status through CDPH. One AO has been granted approval by CDPH and review of the two remaining organizations’ applications is in process. LFS will develop a corrective action plan by December 31, 2015 that includes not only the approval process but also outlines roles and responsibilities for the AOs.

Recommendation 7
Addressing staffing issues by preparing and resubmitting a recruitment and retention proposal, developing a succession plan, and taking necessary steps to implement its planned reorganization.
Response 7
CDPH agrees with this recommendation and is in the process of implementing. LFS has partnered with CDPH’s Human Resources Branch (HRB) to address this recommendation. LFS has loaned two positions to HRB to assist LFS with its recruitment efforts. These positions will allow LFS to work closely with HRB and continue to improve its recruitment, advertising, hiring, and succession planning.

Recommendation 8
Ensuring that its information technology data systems have necessary safeguards, contain accurate and complete data, and support its program needs.

Response 8
CDPH agrees with this recommendation. CDPH’s Information Technology Services Division (ITSD) has already purchased and installed the PEGA Enterprise platform software and has expanded its capabilities to support the many licensing applications within the department. Some of these reusable enhancements that support rapid application development and lower costs include a specialized licensing framework, electronic pay, electronic signature, email communication, CDPH accounting interfaces, and legacy systems interfaces. CDPH has already developed an online personnel licensing system that will be in production by the end of September 2015 which will be the second production application hosted by the PEGA enterprise platform. Also, CDPH is initiating a follow-on project to develop the requirements and functionally required to develop the facilities licensing application on the PEGA Enterprise platform for LFS. In addition, CDPH is actively recruiting staff to support the current and future PEGA Enterprise platform applications.

Recommendation 9
Updating and developing its regulations as necessary to ensure consistency with existing state law.

Response 9
CDPH agrees with this recommendation and has already implemented a plan. As a result of the audit released in 2008, LFS has partnered with CDPH’s Office of Legal Services (OLS) and Office of Regulations to develop a strategy to complete regulations efficiently and timely. This joint effort utilizes a regulations tracking system to identify all of LFS’ regulation packages and to establish timelines for completion of each package. LFS’ packages will update the current regulations to ensure consistency with existing state law. In addition, LFS has committed resources to assist with its regulations efforts.
LFS provided a position to OLS for full-time regulation writing attorney dedicated to LFS. This employee started work in August 2015. In addition, LFS has partnered with OLS to establish an Attorney III position to bolster LFS’ regulations efforts and assist with LFS’ complex needs.