# REPORT TO THE LEGISLATURE

DRUG MANUFACTURING,

MEDICAL DEVICE MANUFACTURING,

AND HOME MEDICAL DEVICE RETAILER

LICENSING COSTS AND FEE ANALYSIS

ANNUAL REPORT

# 2023

CENTER FOR ENVIRONMENTAL HEALTH
DIVISION OF FOOD AND DRUG SAFETY



To obtain a copy of the Division of Food and Drug Safety's Report to the Legislature titled, "Drug Manufacturing, Medical Device Manufacturing, and Home Medical Device Retailer Licensing Costs and Fee Analysis Annual Report 2023," contact:

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# TABLE OF CONTENTS

	EXE	CUTIVE SUMMARYiv
l.	INTE	RODUCTION
	Α.	Background1
	В.	Statutory Requirements for Report to the Legislature2
II.	LICE	ENSE INSPECTION AND INVESTIGATIVE ACTIVITIES
	A.	Drug and Medical Device Manufacturing License
		Inspections and Investigations3
	В.	Home Medical Device Retailer License Inspections and
	Inve	estigations4
III.	LICE	ENSE FEES AND COST ANALYSIS
	Prog	gram License Fees, Revenue, and Program Costs5
		le I – Current Drug and Medical Device Safety Program License Types, entory, and Fees5-6
		le II – Drug and Medical Device Safety Program Projected Revenue by ense Type - Estimated New Applications Fiscal Year (FY) 2023-24
		le III – Drug and Medical Device Safety Program Projected Revenue icense Type - Estimated Renewal Applications FY 2023-246-7
		le IV – Drug and Medical Device Safety Program Operational Costs 2023-248-13

# EXECUTIVE SUMMARY REPORT TO THE LEGISLATURE

## DRUG MANUFACTURING, MEDICAL DEVICE MANUFACTURING, AND HOME MEDICAL DEVICE RETAILER LICENSING COSTS AND FEE ANALYSIS ANNUAL REPORT 2023

California Health and Safety (H&S) Code section 111656.1 (e) requires the California Department of Public Health (CDPH) to annually publish a report to the Legislature recommending the amount of license fees to be charged to drug manufacturers, medical device manufacturers, and home medical device retailer (HMDR) facilities for each Fiscal Year (FY) beginning July 1. H&S Code Section 111656.1 (e) also requires annual license fees to be based on estimated program costs, considering the costs for inspections, investigations, enforcement, and other required activities. The fees collected are deposited into the Drug and Device Safety Fund (Fund 3018) to carry out and implement the licensing provisions of H&S Code, Division 104, Part 5, Chapter 6, Article 6.

The CDPH Food and Drug Branch (FDB) is responsible for licensing all drug and medical device manufacturers in California. Since 1970, CDPH has been mandated to license and inspect drug and medical device manufacturers pursuant to H&S Code section 111635 and to verify products are safe and effective and that firms are operating in compliance with the H&S Code and federal Good Manufacturing Practices (GMP) regulations. Since 2000, FDB has also been responsible for the HMDR Program pursuant to H&S Code section 111656 et seq. Through this program, CDPH licenses and inspects HMDR facilities and warehouses in California that sell and rent home medical devices, and licenses HMDR exemptees (individuals who dispense prescription home medical devices). These inspections verify that the businesses are engaging in sanitary practices so that the devices are being maintained in a manner that protects consumers.

#### I. INTRODUCTION

## A. Background

The Food and Drug Branch (FDB) resides within the California Department of Public Health's (CDPH) Center for Environmental Health, Division of Food and Drug Safety. FDB is responsible for regulating drug manufacturers, medical device manufacturers, home medical device retailer (HMDR) facilities, HMDR warehouses, and HMDR exemptees (an individual that has the knowledge, training, and experience to provide the qualified supervision of the facility), through inspections and maintenance of its licensing program under provisions of the Sherman Food, Drug, and Cosmetic Law (Sherman Law) under Health and Safety (H&S) Code section 109875 et seg. In California, the safety, effectiveness, manufacturing, and labeling of drugs and medical devices have been regulated since the enactment of the California Pure Food and Drugs Act of 1906. Beginning in 1963, drug manufacturers and medical device manufacturers were required to obtain licenses from CDPH before initiating manufacturing operations. Since 1970, CDPH has been mandated to license and inspect drug and medical device manufacturers, pursuant to H&S Code section 111635. Standards for medical devices were not separated from drug standards until 1978 when the federal government promulgated regulations differentiating drugs from medical devices. CDPH began licensing and inspecting HMDR facilities and licensing HMDR exemptees in January 2002, replacing licensing programs by the Board of Pharmacy and the Bureau of Household Furnishing and Thermal Insulation pursuant to Assembly Bill (AB) 1496 (Chapter 837, Statutes of 2000).

CDPH is required to inspect and license drug and medical device manufacturers pursuant to California H&S Code section 111635. The H&S Code adopts federal Good Manufacturing Practices (GMP) regulations for drugs and medical devices (H&S Code section 110105) that establish basic quality assurance standards for manufacturers.

In 1988, a federal law, the Prescription Drug Marketing Act of 1987 (PDMA) (Pub. L. 100-293, 102 Statute 95), was enacted in response to serious public health and safety problems associated with the "diversion market" for prescription drugs. Congress found that adulterated, mislabeled, subpotent, expired, or counterfeit drugs were easily introduced into the national distribution system due to the existence and operation of a wholesale submarket, commonly known as the "diversion market," where drug products are obtained from sources outside of normal channels of distribution. In 1990, the U.S. Food and Drug Administration (FDA) published final regulations establishing state guidelines for the minimum requirements for prescription drug storage and security as well as for the treatment of returned, damaged, and outdated prescription drugs. Further,

wholesale drug distributors were required to establish and maintain inventories and records of all transactions regarding the receipt and distribution of prescription drugs and make these available for inspection and copying by authorized federal, state, or local law enforcement officials. In 1992, California adopted emergency regulations conforming to these FDA requirements, and CDPH began inspecting prescription drug manufacturers for compliance with PDMA regulations.

The licensing inspection requirements for drugs and medical device manufacturers and HMDR facilities are different. Inspections for HMDR facilities are generally uniform and include prelicensing inspections and annual inspections, whereas licensing functions for drug and medical device manufacturers fall into two general types:

- new license applicant evaluation or inspection; or
- for-cause investigations

### B. Statutory Requirements for Report to the Legislature

AB 1496 requires the licensing of HMDR facilities by CDPH. The statute sets specific facility and operational performance standards and requires CDPH to perform inspections prior to licensing each facility. It also requires CDPH to provide an annual report to the Legislature recommending proposed license fee changes based upon the estimated licensing and inspection costs needed to support the program.

#### H&S Code section 111656.1 (e) states:

Commencing January 1, 2003, the department shall, on or before January 10 of each year, provide the Legislature with a report recommending fee rates. The report shall describe the estimated licensing program costs for the next fiscal year to carry out the licensing, regulating, inspecting, and other duties and responsibilities of the department in carrying out the provisions of this article. The department shall describe the projected license fee amount so that license fee revenues cover the estimated licensing program costs. Projected fee amounts shall take into account the resources required for inspections and other activities to support licensing during the previous year and shall take into account projected workload and changes in department overhead costs during the upcoming year.

#### II. LICENSE INSPECTION AND INVESTIGATIVE ACTIVITIES

# A. Drug and Medical Device Manufacturing License Inspections and Investigations

The purpose of the Drug and Medical Device Manufacturing License Program is to prevent the sale and distribution of drug and medical devices that:

- have been improperly manufactured;
- are adulterated, misbranded, or falsely advertised;
- have not been shown to be safe or effective; or
- have not met the medical device design validation requirements as required by the federal quality system regulation

Inspections and investigations of manufacturers allow for the identification and correction of defective products that put California's population at risk. FDB verifies that drug and medical device manufacturers meet GMP requirements and comply with all applicable statutory and regulatory provisions to prevent unqualified and unprepared firms from placing dangerous drugs and medical devices into the hands of medical practitioners and consumers. Firms cannot legally manufacture drugs or medical devices without a valid license. H&S Code section 111635 requires manufacturers to provide CDPH with evidence of ownership and any of the following: valid biologics license, valid FDA registration and inspection, International Organization for Standardization compliance, or an approved investigational new drug or an approved investigational device exemption issued by the FDA.

If the firm is not able to provide the documented evidence as prescribed under H&S Code section 111635, CDPH is required to inspect the place of manufacture prior to issuing a new drug and medical device manufacturing license.

After the initial license is issued, CDPH can inspect or investigate a manufacturer for-cause when CDPH makes the determination that the public health and safety is at risk. Inspections and investigations may additionally be conducted due to complaints, product recalls, or upon request by the FDA to assist with enforcement activities. A license may be denied, suspended, or revoked by CDPH if the manufacturer is found in violation of any applicable part of the Sherman Law.

CDPH drug and medical device manufacturer licensing inspections and investigations verify that:

- new manufacturers have effective quality systems in place before they manufacture and ship drugs and medical devices;
- firms have qualified personnel performing critical process steps;
- all critical drug manufacturing processes, such as purification operations and potency testing, are validated to be appropriate, effective, reproducible, precise, and that firms consistently follow validated processes;
- medical devices are designed and processed with proper validation;
- critical process steps such as sterilization are validated to be effective and that firms consistently follow validated procedures;
- firms investigate all problems that are identified during manufacturing or reported by customers and that corrective and preventive actions are taken to prevent future problems;
- education is provided as needed to help industry understand and comply with manufacturing requirements;
- CDPH has a means to identify when enforcement action is needed to prevent the distribution of unsafe or ineffective drugs; and
- divergent market drug products are not introduced into the national distribution system

### B. Home Medical Device Retailer License Inspections and Investigations

H&S Code section 111656 requires inspection of HMDR facilities prior to licensure and annually thereafter. The licensing of HMDR facilities is important to protect California consumers from unsafe, contaminated medical devices, and adulterated prescription medical oxygen. Mandated license inspections authorize competent and knowledgeable persons (HMDR exemptees) dispense, repair, calibrate, and maintain prescription devices, and sell prescription medical oxygen. The wide variety of medical devices, and how they are assembled, maintained, cleaned, sanitized, and utilized, requires regulatory oversight. Under CDPH's oversight, HMDR facilities distributing or selling medical devices such as ventilators and oxygen concentrators, drug delivery systems for home medication, hospital beds, traction equipment, and other medical devices safely allows consumers to be treated or convalesce at home. Prescription medical oxygen is inspected by CDPH to determine if it is suitable for life support. CDPH coordinates its inspection findings with the California Department of Health Care Services (DHCS) Medi-Cal Provider Enrollment Branch, the Medi-Cal Fraud Prevention Bureau, and Audits and Investigations Division, which includes the Medical Review Branch and the Investigations Branch, to eliminate fraud by identifying HMDR facilities that falsely bill for products never sold or utilized.

#### III. LICENSE FEES AND COST ANALYSIS

### Program License Fees, Revenue, and Program Costs

H&S Code sections 111625, 111630, and 111656.1 authorize CDPH to establish and adjust fees for drug manufacturers, medical device manufacturers, and HMDR licenses. The Drug and Device Safety Fund was established to deposit fees to fund the Drug and Medical Device Safety Program.

The projected revenues for FY 2023-24 are \$6.7 million. The program is projecting total state operations expenses of \$7.9 million, \$278,000 in supplemental pension payments and \$25,000 in Pro Rata for a total of \$8.2 million in estimated expenditures for FY 2023-24. There will not be a fee increase in FY 2023-24. Revenues, in combination with the fund reserve, are estimated to be sufficient to cover expenses for FY 2023-24. Ongoing revenue and expenditure levels will continue to be evaluated in future years for potential fee adjustments.

The Drug and Medical Device Safety Program issues various license types for drug manufacturers, medical device manufacturers, HMDR facilities, and HMDR exemptee licensees under the current licensing fee schedule. Table I provides an existing inventory of current license types with existing fees for FY 2022-23. Table II provides the estimated number of applications, the renewal frequency, and estimated revenue for each type of license for new applicants for FY 2023-24. Table III provides the estimated number of applications, the renewal frequency, and estimated revenue for each type of license for renewal applicants for FY 2023-24.

Table I – Current Drug and Medical Device Safety Program License Types, Inventory, and Fees\*

License Type	Total Inventory*	FY 2022-23 Fee	License Renewal Frequency
Drug Manufacturing License	495	\$4,396	Biennial
Prescription Drug Marketing Act	338	\$200	Biennial
Medical Device Manufacturing License	1386	\$4,396	Biennial
Home Medical Device Retailer	1043	\$1,437	Annual

HDMR Out of State	352	\$254	Annual
HDMR Warehouse	25	\$720	Annual
HDMR Exemptee	2172	New \$423 Renewal \$254	Annual

<sup>\*</sup>Inventory totals are a point-in-time

Table II – Drug and Medical Device Safety Program Projected Revenue by License Type - Estimated New Applications FY 2023-24

Revenue Source (by License Type)	Projected Incoming Applications	FY 2023-24 License Fee	Estimated FY 2023-24 Revenue
Drug Manufacturing License	52	\$4,396	\$228,592
Prescription Drug Marketing Act	41	\$200*	\$8,200
Medical Device Manufacturing License	146	\$4,396	\$641,816
Home Medical Device Retailer	180	\$1,437	\$258,660
HMDR Out of State	88	\$254	\$22,352
HMDR Warehouse	7	\$720	\$5,040
HMDR Exemptee	513	\$423	\$216,999
Total - New	1,045		\$1,381,659

<sup>\*</sup>Licensing fees for Prescription Drug Marketing Act are set in statute; no fee increase implemented.

Table III – Drug and Medical Device Safety Program Projected Revenue by License Type - Estimated Renewal Applications FY 2023-24

Revenue Source (by License Type)	Projected Incoming Renewal Applications	Proposed FY 2023-24 License Fee	Estimated FY 2023-24 Revenue
Drug Manufacturing License	215	\$4,396	\$945,140
Prescription Drug Marketing Act	148	\$200*	\$29,600

Medical Device Manufacturing License	581	\$4,396	\$2,554,076
Home Medical Device Retailer	916	\$1,437	\$1,316,292
HMDR Out of State	269	\$254	\$68,326
HMDR Warehouse	16	\$720	\$11,520
HMDR Exemptee	1625	\$254	\$412,750
Total - Renewal	3,770		\$5,337,704
Total Revenue			\$6,719,363

<sup>\*</sup>Licensing fees for Prescription Drug Marketing Act are set in statute; no fee increase implemented.

To support all licensing and inspection activities, as well as operational activities, the program requires 28.5 staff and associated state operations budget of \$7.9 million in estimated expenditures for FY 2023-24. The number of field and office staff and the cost of the Drug and Medical Device programs are based on the estimated time to complete inspections, compliance investigations, and enforcement activities. In addition to CDPH's requirements to inspect and license drug and medical device manufacturers pursuant to H&S Code section 111635, CDPH is required to follow-up on drug, medical device, and HMDR complaints, alerts, referrals, and recalls. During FY 2021-22, these consisted of 41 additional follow-up activities.

For cause inspections and investigations generally take more time than a new license inspection to complete. These types of investigations are more in-depth and consist of examining corrective actions from the previous inspections, following up on product failures, auditing newly established production processes and resulting records, and evaluating changes and new products. Additional time is necessary to investigate fraudulent activities and develop legal and administrative actions.

Investigators will complete and conduct drug, medical device, and HMDR forcause inspections, complaint referrals, and investigations of a firm's overall compliance with California laws and regulations. Prelicense inspections of drug manufacturers will require an extensive examination of the facility, quality control, employee qualifications, process validation, packaging, labeling, and documentation.

Investigations are conducted by sworn investigators with firms or individuals that are illegally manufacturing or distributing unapproved new drugs or medical

devices. Partnerships with other state and federal regulatory and law enforcement counterparts lead to investigations against illegal importers of unapproved new drug and medical devices. These investigations and enforcement actions can lead to civil or criminal charges or the development of administrative penalty actions. These actions brought on by sworn staff have protected public health and safety, and abated health care fraud.

In addition, program staff review and analyze licensing applications and other required documents, provide technical assistance, provide support to the program, and oversee financial operations. Table IV reflects the operational budget detail for the Drug and Medical Device Safety Program for FY 2023-24.

Table IV - Drug and Medical Device Safety Program Operational Costs FY 2023-24

Position and Function	Salary and Benefits	Operating Expenses	Total cost per FTE	FTE	Total Cost
Chief, Food	201101113	- Apolisos	Politic		10101 0001
and Drug	\$326,788	\$36,000	\$362,788	1.0	\$362,788
Section	second-level personnel; resactivities specificensing progractivities assuming lement the compliant firm reports, and coordination consistency of their activities programs. Defined deviation deviation and the consistency of their activities programs. Defined deviation deviation of the consistency of their activities programs. Defined deviation d	supervisory se sponsible for in cific to the dru gram. Coordin ring resources e program an ns. Prepare pe oversee staff of of scientific a of investigative s with other di velop, mainto ce, and HMDI ning, education assure product input to regulated thers related to PH at relevan	ntific, administration staff of law mplementing are propertied and expertise and expertise and expertise and expertise and expertise and development. On the control of the	w enforce and monited device and engand ensed and ensed and ensed and ensed and ensed and ensed	ement oring the units and its nforcement able to nd non- probation and direct the o assure rcement, and e and federal vide drug, s, procedures, ctivities technical and effectiveness nsultation to a regarding vices;

Position and Function	Salary and Benefits	Operating Expenses	Total cost per FTE	FTE	Total Cost
Staff Services	Dericinis	EXPENSES	PCITIE		
Manager I	\$159,901	\$26,000	\$185,901	0.5	\$92,951
	· ·	sis of progran	n data collecte	d. Provid	
	-		vel of program		
	•		nance of compl		
	•		acking system fo		
			nsing medical d		
			ure requirement		
		•	products. Analy	. •	
		_	orts. Oversee ar		
			nment of licensir edures for new	_	_
	-	-	d administrative		_
			ants and issue o	•	
		• •	n budget to mo		
			he revenue mo		
	of the fund. R	espond to co	mplex issues rela	ated to in	nventory,
	•		e Program. Prep	oare repo	orts for
	managemen	t. -	T	T	I
Investigator	\$204,765	\$36,000	\$240,765	6.0	\$1,444,590
	Train and lead	d staff in the p	erformance of	field inve	estigations and
			x criminal, civil, (		
			verify suspected		
			riminal/civil inve	_	
		•	cted of major he		
	•	_	on includes gatl	_	
	prescribed by law; interviewing/interrogating witnesses and				
	suspects; developing investigation plans; developing operational plans and safely executing them; conducting surveillance and				
	covert activities; preparing detailed reports and making				
	recommendations based on state and federal laws/regulations;				
	preparing and executing search warrants, inspection warrants,				
	arrest warrants and subpoenas, and testifying in court or				
	administrative proceedings. Prevent unapproved new drugs and				
	medical devices from being received into commerce and/or from being sold outside the normal distribution chain.				
	trom being so	id outside the	normal distribu	tion cha	ın.

Position and Function	Salary and Benefits	Operating Expenses	Total cost per FTE	FTE	Total Cost
Environmental Scientist	\$129,842	\$36,000	\$165,842	11.0	\$1,824,262
	Inspect drug and medical device manufacturers and HMDR facilities. Perform preliminary and secondary analysis, research, and surveys of typical drug and medical device manufacturing and HMDR practices. Prepare for inspection activities and complete license inspections to determine ownership, adequacy of facilities, personnel qualifications, and compliance with applicable regulations.				
Senior Environmental	\$156,047	\$26,000	\$182,047	2	\$364,094
Scientist (Specialist)	contract inspections of the contract status investigations. HMDR practic Assist Environment inspection regard conduct Provide Environment of the conduct of	ections, review us. Provide tra of drug and reces. Prepare E mental Scienti ports. Audit/Recents the most high conmental Scies sults of prelimi reys, and inve	conducting FDA we reports, track, ining related to medical device nvironmental So st Supervisor with eview employed and technical and the reconstigations of typopractices.	and mo surveys of manufa cientist w h evaluce e perform d compl g. Assist i	nitor the and cturing and ork plans. Ition of hance. Lead ex inspections. In the alysis,
Senior Environmental	\$233,783	\$26,000	\$259,783	2.0	\$519,566
Scientist (Supervisory)	Provide super investigations HMDR practic assignments. employee per Development Supervise and inspections. Provide supervise s	vision and tra of drug and reseas. Prepare E Evaluate insper rformance. Profermance Profermance A definition of the rovide Enviror of the sof preliming nvestigations	ining related to medical device nvironmental Scetion reports. A repare Individuo ppraisal Plans a most highly technental Scientisary and secondof typical medical	surveys of manufa cientist w Audit/Reval nd proba chnical a t field tro ary analy	cturing and ork view ation reports. nd complex sining.

Position and	Salary and	Operating	Total cost		
Function	Benefits	Expenses	per FTE	FTE	Total Cost
Supervising Food and	\$261,377	\$36,000	\$297,377	1.0	\$297,377
Drug	•	•	ining, evaluate	l	
Investigator	The second secon		estigations and	-	
	7		nts. Evaluate ins		-
	Audit/Review	employee pe	erformance. Pre	pare Ind	ividual
			ppraisal Plans a	-	=
			ntry. Provide inv	_	_
		•	ince investigation	•	
	_		olete complex c Conduct hearin	•	
	enforcement	•	soriadei riediiri	gs and o	711101
Food and					
Drug Program	\$265,209	\$36,000	\$301,209	2.0	\$602,418
Specialist			acilitate investi	•	-
			ampling, and er		
		_	oals and objecti	=	_
	•	•	ngths and defic ninistrative and		
		-	gram deficienc		
	evaluate mor	-	gram denom	7,03. 1.0 11	ovv arra
		,	etermine pattei	rns or trei	nds in the
	drug, and me	dical device	manufacturing	industry.	Coordinate
			cts and survey a		
		•	omplex FDA refe		_
		•	check quality co perience, and iss		
	1	•	expert on drug		
			issues. Develop		
		•	explain laws, reg	•	
	enforcement	policies.			

Position and	Salary and	Operating	Total cost			
Function	Benefits	Expenses	per FTE	FTE	Total Cost	
Staff Services	4107100	401000	4100100		<b>*</b> 100.100	
Analyst	\$107,132   \$26,000   \$133,132   1.0   \$133,132   Oversee financial operations and provide program support. Review					
		•				
			ations and other r le for information	•		
		_	with new applica			
	,	•	d/or other outsta		,	
	identified by t	he investigators	via written corre	spondenc	ce. Verify	
	_	•	mpliance with sto	•	· ·	
	•	_	a and knowledge			
	•		ce with Unit proc			
		_	ial information fro atements submit		•	
	manufacturer			rod by di	09	
Associate				1		
Associate	¢100 041	000	¢1 <i>EE</i> 0 <i>A</i> 1	1.0	¢155 0 / 1	
Governmenta I Program	\$129,841	\$26,000	\$155,841 a variety of tas	1.0	\$155,841	
Analyst		•	and complex, a		_	
7 (TIGIYSI		•	Compliance an	-		
	_		Device/Cosmeti			
	` '	•	uties such a dev		•	
	~	-	rocedures and o			
		• • • • • • • • • • • • • • • • • • • •	Sherman Food			
	Laws Act and	d the Board of	Pharmacy Law	to make	informed	
	decisions and	d align busines	ss practices for [	OMDCES.	Responding	
			ne public, exterr			
			nsing requireme			
	-	•	ility for all MDSU	-	·	
			team leader a		dinate the	
Chiof Food	ettorts of mul	iliaiscipiinary a	rug and device	reams.		
Chief, Food and Drug Unit	\$302,993	\$36,000	\$338,993	1.0	\$338,993	
drid brog orin						
	Second line supervisor of regulatory and law enforcement personnel. Manage program staff performing activities specific					
	to the drug and medical device units and its licensing program.					
	Coordinate the licensing activities assuring resources are					
	available to implement the licensing program and identify					
		•	liant firms. Prepo		-	
	evaluations,	probation rep	orts, and overse	e staff d	evelopment.	
	Oversee the	legal, investige	ative, educatior	nal and e	enforcement	

activities specific to the drug and medical device units. Track and forecast emerging public health trends affecting the licensed industries, and develop regulatory strategies to address them. Develop and direct staff specifically to respond to: 1) adulterated, misbranded, falsely advertised, or otherwise unsafe drugs and medical devices; 2) unsafe or otherwise improper production and processing practices; and 3) drug and medical device recalls and complaint investigations. Assure utilization of state-of-the-art scientific and technical developments that can detect and prevent consumers from being exposed to unsafe products and production practices. Evaluate and audit statewide licensing program plans, policies, procedures, budgets, training, education efforts, and all other activities necessary to assure product safety. Collaborate with the FDA, Board of Pharmacy, Medical Board of California, and other regulatory agencies to develop work plans and share issues and concerns to protect public health. Oversee the scientific, legal, investigative, educational, and enforcement activities specific to the work of their respective unit.

Total Staff Cost		\$6,136,012
CDPH Distributed and Administrative Costs		\$1,736,491
Total Program Operations Estimated Cost	28.5	\$7,872,503