

[Health Code - Regulating Medical Specimen Test Collection Sites]

Ordinance amending the Health Code to require that sites that collect medical specimens on behalf of clinical laboratories partner with either a governmental entity, a licensed health care provider located in the City, or an educational or academic institution, establish hygiene, sanitation, and privacy standards, and adhere to the Health Insurance Portability and Accountability Act; prohibiting such sites from paying individuals to take a medical test; and providing that a violation of the specimen collection standards is a misdemeanor offense and a public health nuisance subject to an administrative penalty that may be imposed by the Department of Public Health.

NOTE: Unchanged Code text and uncodified text are in plain Arial font. Additions to Codes are in *single-underline italics Times New Roman font*. Deletions to Codes are in *strikethrough italics Times New Roman font*. Board amendment additions are in double-underlined Arial font. Board amendment deletions are in ~~Arial font~~. Asterisks (\* \* \* \*) indicate the omission of unchanged Code subsections or parts of tables.

Be it ordained by the People of the City and County of San Francisco:

Section 1. The Health Code is hereby amended by adding Article 49, consisting of Sections 4901 through 4906, to read as follows:

**ARTICLE 49:**

**SPECIMEN TEST COLLECTION SITES**

**SEC. 4901. FINDINGS AND PURPOSE.**

(a) Since the onset of the COVID-19 emergency, it has become increasingly common for City residents to see organizations and businesses operate clinical testing sites on City sidewalks and in

1 other public locations. Medical testing sites that both collect specimens and then perform clinical tests  
2 on those specimens are called “laboratories” or “clinical laboratories,” and are licensed and  
3 regulated by federal Centers for Medicare and Medicaid Services and the California Department of  
4 Public Health or the applicable state agency for laboratories outside California. By contrast, sites that  
5 collect specimens but do not actually perform clinical tests, and merely send the specimens to a  
6 laboratory for testing (“Specimen Collection Sites”) are not subject to CMS or CDPH regulation or  
7 oversight.

8 (b) Generally, under the Health Insurance Portability and Accountability Act and its  
9 implementing regulations (collectively, “HIPAA”), Specimen Collection Sites that collect specimens on  
10 behalf of covered entities, such as clinical laboratories, are business associates of those covered  
11 entities as those terms are defined under HIPAA. Business associates are obligated to follow HIPAA’s  
12 privacy and safety requirements.

13 (c) It is critically important that Specimen Collection Sites in San Francisco protect the privacy  
14 of individuals’ health information and comply with health and safety protocols for handling and testing  
15 infectious disease specimens. Accordingly, the purpose of this Article 49 is to set forth the minimum  
16 privacy and health and safety requirements for Specimen Collection Sites to ensure that such sites are  
17 operating in a safe and lawful manner.

18  
19 **SEC. 4902. DEFINITIONS.**

20 For purposes of this Article 49, the following terms have the following meanings:

21 “CDC” means the federal Centers for Disease Control and Prevention.

22 “CDPH” means the California Department of Public Health.

23 “City” means the City and County of San Francisco.

1           “CLIA” means the Clinical Laboratory Improvement Amendments, codified at 42 U.S.C.  
2 § 263a, as it may be amended from time to time, and including any implementing regulations or  
3 guidance promulgated by CMS, the CDC, or the federal Food and Drug Administration.

4           “CMS” means the federal Centers for Medicare and Medicaid Services.

5           “Covered Operator” means a private, for-profit or non-profit person, company, or other  
6 organization operating one or more Specimen Collection Sites anywhere in the City. Covered  
7 Operator includes a person, company, or organization that collects specimens without charge to the  
8 Examinee, regardless of whether reimbursement or payment is sought from insurance companies or  
9 federal, state, or local governmental agencies. Covered Operator does not include government entities  
10 or any facility (such as a general acute care hospital, skilled nursing facility, or ambulatory clinic) that  
11 directly collects specimens and is subject to regulation by CDPH.

12           “Department” means the San Francisco Department of Public Health.

13           “Director” means the Director of Health, or the Director’s designee.

14           “Examinee” means an individual providing a specimen to the Specimen Collection Site.

15           “Personnel” means employees, contractors and sub-contractors, including but not limited to  
16 those who sell goods or perform services onsite or who deliver goods for the Covered Operator,  
17 vendors who are permitted to sell goods onsite, volunteers, and other individuals who regularly provide  
18 services to a Covered Operator.

19           “PPE” means Personal Protective Equipment.

20           “Specimen Collection Site” means a site where a Covered Operator or its Personnel obtain  
21 specimens for testing for medical or health conditions, including by way of example but not limitation,  
22 COVID-19 and flu, from an Examinee and for delivery to an off-site CLIA-certified laboratory for  
23 clinical processing. Specimen Collection Sites do not include sites regulated by CMS or CDPH where  
24 clinical laboratory tests are performed on the premises.

1 “Test” means the diagnostic test used to detect any infectious, contagious, or  
2 communicable disease and that the Covered Operator sends to a CLIA-certified laboratory for  
3 clinical processing.

4 “Well-Fitted Mask” means a face covering that is well-fitted to an individual and covers the  
5 nose and mouth while talking. A Well-Fitted Mask does not include a scarf, ski mask, balaclava,  
6 bandana, turtleneck, collar, or single layer of fabric, or any mask that has an unfiltered one-way  
7 exhaust valve.

8  
9 **SEC. 4903. REQUIREMENTS FOR SPECIMEN COLLECTION SITES.**

10 (a) Each Covered Operator must provide its Personnel with guidelines for wearing appropriate  
11 PPE based on the type of specimen to be collected by Personnel. Covered Operators must provide  
12 Personnel with information and training on the proper procedures for putting on and taking off PPE  
13 based on the type of specimen collected by the Personnel. Each Covered Operator’s guidelines for  
14 wearing appropriate PPE must include the following minimum standards:

15 (1) If collecting specimens or working within six feet of Examinees, Personnel must  
16 wear a Well-Fitted Mask, eye protection, gloves, and a gown.

17 (2) Personnel who handle specimens, but are not directly involved in collection (e.g.,  
18 handling self-collected specimens) and not working within six feet of the Examinee, must wear a Well-  
19 Fitted Mask and gloves.

20 (3) Personnel must change gloves after handling a specimen or whenever their gloves  
21 become soiled or torn.

22 (b) Personnel at Specimen Collection Sites must designate a surface area for specimen  
23 collection and handling and disinfect that area using a disinfectant product registered with the federal  
24 Environmental Protection Agency for use against contagious, infectious, or communicable diseases.  
25 Personnel must disinfect the surface areas at the following times: (1) before specimen collection begins

1 each day; (2) after Personnel collect a specimen; (3) when visibly soiled; (4) in the event of a specimen  
2 spill; and (5) at the end of every day. Each Covered Operator must at all times during hours of  
3 operation make hand sanitizer available for use by Personnel and Examinees.

4 (c) Each Covered Operator must provide all Examinees a written informed consent form  
5 consenting to the collection of the specimen and the testing of that specimen. Before the specimen is  
6 collected, the Examinee must sign the informed consent form. Personnel must provide a copy of the  
7 signed form, either in hard copy or electronically, to the Examinee.

8 (d) Each Covered Operator must have written policies covering the following topics:

9 \_\_\_\_\_ (1) Specimen collection, storage, and transport, that addresses the specific types of  
10 specimens the Specimen Collection Site will collect or are consistent with the test manufacturers'  
11 instructions.

12 \_\_\_\_\_ (2) Training of Personnel in PPE requirements; specimen collection, storage, and  
13 transport; and protection of personal information of Examinees seeking or considering seeking medical  
14 testing at the Specimen Collection Site.

15 \_\_\_\_\_ (3) Test Result Notification, including how results are provided to Examinees either by  
16 the Covered Operator, its Personnel, or by the CLIA-certified laboratory where the specimens are  
17 tested.

18 \_\_\_\_\_ (4) A privacy policy regarding Examinees' medical and health information, biological  
19 samples, and test results.

20 The written policies and procedures specified in subsection (d)(1) through (d)(4) must be  
21 provided to: all Personnel; any member of the public, upon request, including, but not limited to,  
22 Examinees seeking or considering seeking medical testing at a Specimen Collection Site; City, state, or  
23 federal employees conducting inspections or investigations; and any CLIA-certified laboratory where  
24 the specimens will be tested to enable the lab to verify the integrity of the specimens being collected.

1 (e) Covered Operators may use human biological/viral specimens only for (1) clinical testing  
2 and (2) laboratory validation and quality control, to the extent such uses are allowed by applicable  
3 laws, rules, regulations, and licensure requirements.

4 (f) Upon request by any member of the public, including, but not limited to, Examinees seeking  
5 or considering seeking testing at a Specimen Collection Site, and City, state, or federal employees  
6 conducting inspections or investigations, Personnel at Specimen Collection Sites must produce the  
7 name of the Specimen Collection Site's ordering/prescribing provider, where a prescription is required  
8 for collection of samples and processing by CDPH-approved laboratories.

9 (g) Upon request by any member of the public, including, but not limited to, Examinees seeking  
10 or considering seeking testing at a Specimen Collection Site, and City, state, or federal employees  
11 conducting inspections or investigations, Personnel at Specimen Collection Sites must produce the  
12 following documentation from the laboratory that will be processing/performing Tests on the  
13 specimens collected at the Specimen Collection Site: (1) a current and valid CLIA license; and (2) a  
14 current and valid Clinical and Public Health Laboratory License from CDPH.

15 (h) A Specimen Collection Site operated by a Covered Operator must comply with all  
16 applicable privacy laws, including but not limited to HIPAA. In the event HIPAA does not apply to the  
17 Covered Operator, then the Covered Operator must adhere to the same standards as provided by  
18 HIPAA to safeguard Examinee confidentiality and medical information.

19 (i) Each Covered Operator must partner with one of the following entities to perform Tests on  
20 behalf of the entity: (a) a governmental entity; (b) a licensed health care provider located in the City;  
21 or (c) an educational or academic institution (including but not limited to licensed child care providers,  
22 preschools, public and private schools, colleges, universities, and similar institutions of higher  
23 learning). Upon request, Personnel at a Specimen Collection Site must demonstrate evidence  
24 of the partnership with one of the foregoing entities by producing a written agreement,  
25 memorandum, letter, or similar document that shows the entity has requested the Specimen

1 Collection Site perform Tests on behalf of the entity. The ordering prescriber's standing order,  
2 required by subsection (f), shall not constitute sufficient evidence of a partnership.

3 (j) Covered Operators shall not offer or pay Examinees any remuneration, including  
4 anything of value, directly or indirectly, overtly or covertly, in cash or in kind, to induce, or in  
5 exchange for, the Examinee using a Test offered by the Covered Operator. The prohibition  
6 on remuneration shall not apply to clinical testing performed pursuant to an institutional review  
7 board-approved research study or with the approval of the Department.

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9 **SEC. 4904. ADMINISTRATION AND ENFORCEMENT.**

10 (a) This Article 49 shall be administered and enforced by the Department. The Director may  
11 adopt regulations, guidelines, and forms to carry out the provisions and purposes of this Article 49.

12 (b) For purposes of assessing penalties for violation of Section 4903, each instance that a  
13 Specimen Collection Site violates any provision of Section 4903 shall constitute a separate violation.

14 (c) The Director may issue a notice of violation for violations of Section 4903. The Director  
15 may impose an administrative penalty of not less than \$250 and not more than \$1,000 per violation.  
16 Administrative Code Chapter 100, "Procedures Governing the Imposition of Administrative Fines," is  
17 hereby incorporated in its entirety, except: (1) as it relates to the definition of a violation and the  
18 calculation of penalty amounts, addressed in Sections 4904(b) and (c); and (2) that the Director shall  
19 appoint the hearing officer to conduct hearings for appeals.

20 (d) A violation of Section 4903 shall be considered a nuisance under Health Code Section 581,  
21 or any successor provision.

22 (e) The Department shall have authority to enforce Section 4903 under Health Code Sections  
23 594, 595, 596, 596.5, 599, 600, and 610.

24  
25 **SEC. 4905. VIOLATION A MISDEMEANOR.**

1           Any person who violates Section 4903 is guilty of a misdemeanor. Any person  
2 convicted of a misdemeanor hereunder is punishable by a fine of not more than \$500 or by  
3 imprisonment for a period of not more than six months, or by both. A person who violates the  
4 provisions of Section 4903 is guilty of a separate offense for each day, or portion thereof,  
5 during which the violation continues.

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7           **SEC. 49065. UNDERTAKING FOR THE GENERAL WELFARE.**

8           In enacting and implementing this Article 49, the City is assuming an undertaking only to  
9 promote the general welfare. It is not assuming, nor is it imposing on its officers and employees, an  
10 obligation for breach of which it is liable in money damages to any person who claims that such breach  
11 proximately caused injury.

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13           **SEC. 49076. SEVERABILITY.**

14           If any section, subsection, sentence, clause, phrase, or word of this Article 49, or any  
15 application thereof to any person or circumstance, is held to be invalid or unconstitutional by a  
16 decision of a court of competent jurisdiction, such decision shall not affect the validity of the remaining  
17 portions or applications of this Article. The Board of Supervisors hereby declares that it would have  
18 passed this Article and each and every section, subsection, sentence, clause, phrase, and word not  
19 declared invalid or unconstitutional without regard to whether any other portion of this Article or  
20 application thereof would be subsequently declared invalid or unconstitutional.

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22           Section 2. Effective Date. This ordinance shall become effective 30 days after  
23 enactment. Enactment occurs when the Mayor signs the ordinance, the Mayor returns the  
24 ///







**City and County of San Francisco**  
**Tails**  
**Ordinance**

City Hall  
1 Dr. Carlton B. Goodlett Place  
San Francisco, CA 94102-4689

**File Number:** 231158

**Date Passed:** January 30, 2024

Ordinance amending the Health Code to require that sites that collect medical specimens on behalf of clinical laboratories partner with either a governmental entity, a licensed health care provider located in the City, or an educational or academic institution, establish hygiene, sanitation, and privacy standards, and adhere to the Health Insurance Portability and Accountability Act; prohibiting such sites from paying individuals to take a medical test; and providing that a violation of the specimen collection standards is a misdemeanor offense and a public health nuisance subject to an administrative penalty that may be imposed by the Department of Public Health.

November 07, 2023 Board of Supervisors - AMENDED, AN AMENDMENT OF THE WHOLE BEARING NEW TITLE

Ayes: 9 - Chan, Dorsey, Engardio, Mandelman, Peskin, Preston, Safai, Stefani and Walton

Excused: 2 - Melgar and Ronen

November 07, 2023 Board of Supervisors - RE-REFERRED AS AMENDED

Ayes: 9 - Chan, Dorsey, Engardio, Mandelman, Peskin, Preston, Safai, Stefani and Walton

Excused: 2 - Melgar and Ronen

January 11, 2024 Public Safety and Neighborhood Services Committee - RECOMMENDED

January 23, 2024 Board of Supervisors - PASSED ON FIRST READING

Ayes: 11 - Chan, Dorsey, Engardio, Mandelman, Melgar, Peskin, Preston, Ronen, Safai, Stefani and Walton

January 30, 2024 Board of Supervisors - FINALLY PASSED

Ayes: 11 - Chan, Dorsey, Engardio, Mandelman, Melgar, Peskin, Preston, Ronen, Safai, Stefani and Walton

File No. 231158

I hereby certify that the foregoing Ordinance was FINALLY PASSED on 1/30/2024 by the Board of Supervisors of the City and County of San Francisco.



Angela Calvillo  
Clerk of the Board



London N. Breed  
Mayor

2/9/24

Date Approved