

Kroll Laboratory Specialists, Inc., Scientific Testing Laboratories, Inc.; Kroll Scientific Testing Laboratories, Inc.)

Clinical Reference Laboratory, Inc., 8433 Quivira Road, Lenexa, KS 66215–2802, 800–445–6917

Cordant Health Solutions, 2617 East L Street, Tacoma, WA 98421, 800–442–0438 (Formerly: STERLING Reference Laboratories)

Desert Tox, LLC, 5425 E Bell Rd., Suite 125, Scottsdale, AZ 85254, 602–457–5411/623–748–5045

DrugScan, Inc., 200 Precision Road, Suite 200, Horsham, PA 19044, 800–235–4890

Dynacare, * 245 Pall Mall Street, London, ONT, Canada N6A 1P4, 519–679–1630 (Formerly: Gamma-Dynacare Medical Laboratories)

Upon finding a Canadian laboratory to be qualified, HHS will recommend that DOT certify the laboratory (**Federal Register**, July 16, 1996) as meeting the minimum standards of the Mandatory Guidelines published in the **Federal Register** on January 23, 2017 (82 FR 7920). After receiving DOT certification, the laboratory will be included in the monthly list of HHS-certified laboratories and participate in the NLCP certification maintenance program.

ElSohly Laboratories, Inc., 5 Industrial Park Drive, Oxford, MS 38655, 662–236–2609

Laboratory Corporation of America Holdings, 7207 N. Gessner Road, Houston, TX 77040, 713–856–8288/800–800–2387

Laboratory Corporation of America Holdings, 69 First Ave., Raritan, NJ 08869, 908–526–2400/800–437–4986, (Formerly: Roche Biomedical Laboratories, Inc.)

Laboratory Corporation of America Holdings, 1904 TW Alexander Drive, Research Triangle Park, NC 27709, 919–572–6900/800–833–3984, (Formerly: LabCorp Occupational Testing Services, Inc.; CompuChem Laboratories, Inc.; CompuChem Laboratories, Inc., A Subsidiary of

Roche Biomedical Laboratory; Roche CompuChem Laboratories, Inc., A Member of the Roche Group)

Laboratory Corporation of America Holdings, 1120 Main Street, Southaven, MS 38671, 866–827–8042/800–233–6339, (Formerly: LabCorp Occupational Testing Services, Inc.; MedExpress/National Laboratory Center)

LabOne, Inc. d/b/a Quest Diagnostics, 10101 Renner Blvd., Lenexa, KS 66219, 913–888–3927/800–873–8845, (Formerly: Quest Diagnostics Incorporated; LabOne, Inc.; Center for Laboratory Services, a Division of LabOne, Inc.)

Legacy Laboratory Services Toxicology, 1225 NE 2nd Ave., Portland, OR 97232, 503–413–5295/800–950–5295,

MedTox Laboratories, Inc., 402 W. County Road D, St. Paul, MN 55112, 651–636–7466/800–832–3244

Minneapolis Veterans Affairs Medical Center, Forensic Toxicology Laboratory, 1 Veterans Drive, Minneapolis, MN 55417, 612–725–2088, Testing for Veterans Affairs (VA) Employees Only.

Pacific Toxicology Laboratories, 9348 DeSoto Ave., Chatsworth, CA 91311, 800–328–6942, (Formerly: Centinela Hospital Airport Toxicology Laboratory)

Pathology Associates Medical Laboratories, 110 West Cliff Dr., Spokane, WA 99204, 509–755–8991/800–541–7891x7

Phamatech, Inc., 15175 Innovation Drive, San Diego, CA 92128, 888–635–5840

Quest Diagnostics Incorporated, 1777 Montreal Circle, Tucker, GA 30084, 800–729–6432, (Formerly: SmithKline Beecham Clinical Laboratories; SmithKline Bio-Science Laboratories)

Quest Diagnostics Incorporated, 400 Egypt Road, Norristown, PA 19403, 610–631–4600/877–642–2216, (Formerly: SmithKline Beecham Clinical Laboratories; SmithKline Bio-Science Laboratories)

Redwood Toxicology Laboratory, 3700 Westwind Blvd., Santa Rosa, CA 95403, 800–255–2159,

U.S. Army Forensic Toxicology Drug Testing Laboratory, 2490 Wilson St., Fort George G. Meade, MD 20755–5235, 301–677–7085, Testing for Department of Defense (DoD) Employees Only

Anastasia Marie Donovan,
Policy Analyst.

[FR Doc. 2020–14040 Filed 6–29–20; 8:45 am]

BILLING CODE 4162–20–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Substance Abuse and Mental Health Services Administration

Agency Information Collection Activities: Submission for OMB Review; Comment Request

Periodically, the Substance Abuse and Mental Health Services Administration (SAMHSA) will publish a summary of information collection requests under OMB review, in compliance with the Paperwork Reduction Act (44 U.S.C. Chapter 35). To request a copy of these documents, call the SAMHSA Reports Clearance Officer on (240) 276–0361.

Project: Mandatory Guidelines for Federal Workplace Drug Testing Programs (OMB No. 0930–0158)—Revision

SAMHSA will request OMB approval for a revision of the Federal Drug Testing Custody and Control Form (CCF) for Federal agency and federally regulated drug testing programs which must comply with the HHS Mandatory Guidelines for Federal Workplace Drug Testing Programs using Urine (UrMG) dated January 23, 2017 (82 FR 7920) and using Oral Fluid (OFMG) dated October 25, 2019, and OMB approval for information provided by test facilities (laboratories and Instrumented Initial Test Facilities, IITFs) for the National Laboratory Certification Program (NLCP).

The CCF is used by all Federal agencies and employers regulated by the Department of Transportation (DOT) and the Nuclear Regulatory Commission (NRC) to document the collection and chain of custody of urine specimens at the collection site, for HHS-certified test facilities to report results, and for Medical Review Officers (MROs) to document and report a verified result. SAMHSA allows the use of the CCF as a paper or electronic form.

The current OMB-approved CCF has an August 31, 2020 expiration date. SAMHSA has resubmitted the CCF with revisions to the form for OMB approval. During 60-day public comment 7 commenter's submitted comments on the proposed changes to the CCF. These commenters were comprised of individuals, organizations, and private sector companies. All comments were reviewed and taken into consideration in the preparation of the revised CCF. The issues and concerns raised in the public comments for the CCF are set out www.reginfo.gov/public/do/PRAMain.

These revisions are listed below:

* The Standards Council of Canada (SCC) voted to end its Laboratory Accreditation Program for Substance Abuse (LAPSA) effective May 12, 1998. Laboratories certified through that program were accredited to conduct forensic urine drug testing as required by U.S. Department of Transportation (DOT) regulations. As of that date, the certification of those accredited Canadian laboratories will continue under DOT authority. The responsibility for conducting quarterly performance testing plus periodic on-site inspections of those LAPSA-accredited laboratories was transferred to the U.S. HHS, with the HHS' NLCP contractor continuing to have an active role in the performance testing and laboratory inspection processes. Other Canadian laboratories wishing to be considered for the NLCP may apply directly to the NLCP contractor just as U.S. laboratories do.

Copies 1–5

Revised Step 1

1. Added “CDL State and No.” to donor identification types
2. Added “Collector Contact Info:” and “Other” line (e.g., email)

Revised Step 2

1. Put Urine and Oral Fluid checkboxes above Step 2 for collector to annotate
2. Expanded to 4 lines for collector entries:
 - General entry for Split, Single, or None Provided (same as current)
 - Entries specific to urine collection (moved “Collector reads urine temperature within 4 minutes” here; other entries same as current)
 - Entries specific to oral fluid collection: Added “Split Type” with checkboxes for Serial, Concurrent, and Subdivided; “Each Device Within Expiration Date?” with checkboxes Yes or No; and Volume Indicator(s) Observed checkbox)
 - Remarks (same as current)

Revised Step 3

1. Edited instruction to state “collector affixes seal(s) to bottle(s)/tube(s)”

Revised Step 4 (Collector Section)

1. Edited “Specimen Bottle(s) Released To” box to state “Specimen Bottle(s)/Tubes(s) Released To”

Copy 1 (Test Facility Copy)

Revised Step 4 (Accessioner Section)

1. Edited “Specimen Bottle(s) Released To” box to state “Specimen Bottle(s)/Tubes(s) Released To”

2. Added “Primary/Single Specimen Device Expiration Date” and “Split Specimen Device Expiration Date” fields for accessioner to annotate expiration dates of oral fluid collection devices

Revised Step 5a (Certification and Reporting Section)

1. Removed analyte names and checkboxes
2. Repositioned results and checkboxes: Moved REJECTED FOR TESTING, ADULTERATED, SUBSTITUTED and INVALID RESULT checkboxes; moved POSITIVE checkbox to be under DILUTE
3. Added line for certifying scientist to record positive analytes and concentrations, and added “Analyte(s) in ng/mL” instruction (aligned under “POSITIVE for:”)

Copy 2 (Medical Review Officer Copy)

Revised Step 5 (Donor Section)

1. Added line for donor email address
2. Edited donor certification statement to state “specimen bottle/tubes”

Revised Step 6 (MRO section—Primary Specimen)

1. Put Urine and Oral Fluid checkboxes above Step 6 for MRO to annotate

Bottom of Copies

Revised Copy 1

1. Edited label/seal at bottom of Copy 1 to allow for modification (e.g., perforations, label with transparent seal on one side, and separate label and seal)

Revised Copy 5

1. Removed Instructions for Completing the CCF from the back. SAMHSA will post instructions for completing the Federal CCF for urine and oral fluid on their website.

Based upon information from Federal agencies and from DOT concerning their regulated industries, the number of respondents has increased from 5.4 million to 6.7 million, which increases the total burden hours by 170,701.8–C.;

Laboratories and IITFs seeking HHS certification under the NLCP must complete and submit the NLCP application form. The NLCP application form has not been revised compared to the previous form.

Prior to an inspection, an HHS-certified laboratory or IITF is required to submit specific information regarding its procedures. Collecting this information prior to an inspection allows the inspectors to thoroughly review and understand the testing procedures before arriving for the onsite inspection. The NLCP information checklist has not been revised compared to the previous form.

The annual total burden estimates for the CCF, the NLCP application, the NLCP information checklist, and the NLCP recordkeeping requirements are shown in the following table.

Form/respondent	Number of respondents	Responses per respondent	Total number of responses	Burden per response (hours)	Annual burden (hours)
Custody and Control Form: 1					
Donor	6,726,610	1	6,726,610	0.08	538,128.8
Collector	6,726,610	1	6,726,610	0.07	378,000
Laboratory	6,726,610	1	6,726,610	0.05	336,330
IITF	1	0	0	0.05	0
Medical Review Officer	6,726,610	1	6,726,610	0.05	270,000
NLCP Application Form: 2					
Laboratory	5	5	5	3	15
IITF	0	0	0	3	0
Sections B and C—NLCP Inspection Checklist:					
Laboratory	29	1	29	1	29
IITF	0	0	0	1	0
Record Keeping:					
Laboratory	29	1	29	250	7,250
IITF	0	0	0	250	0
Total	6,726,673	26,906,503	1,529,753

Written comments and recommendations for the proposed information collection should be sent

within 30 days of publication of this notice to www.reginfo.gov/public/do/PRAMain. Find this particular

information collection by selecting “Currently under 30-day Review—Open

for Public Comments” or by using the search function.

Carlos Graham,

Social Science Analyst.

[FR Doc. 2020–13986 Filed 6–29–20; 8:45 am]

BILLING CODE 4162–20–P

DEPARTMENT OF HOMELAND SECURITY

U.S. Customs and Border Protection

[1651–0131]

Agency Information Collection Activities: e-Allegations Submission

AGENCY: U.S. Customs and Border Protection (CBP), Department of Homeland Security.

ACTION: 60-Day notice and request for comments; extension of an existing collection of information.

SUMMARY: The Department of Homeland Security, U.S. Customs and Border Protection will be submitting the following information collection request to the Office of Management and Budget (OMB) for review and approval in accordance with the Paperwork Reduction Act of 1995. The information collection is published in the **Federal Register** to obtain comments from the public and affected agencies.

DATES: Comments are encouraged and will be accepted no later than August 31, 2020 to be assured of consideration.

ADDRESSES: Written comments and/or suggestions regarding the item(s) contained in this notice must include the OMB Control Number 1651–0131 in the subject line and the agency name. To avoid duplicate submissions, please use only *one* of the following methods to submit comments:

(1) Email. Submit comments to: CBP_PRA@cbp.dhs.gov.

(2) Mail. Submit written comments to CBP Paperwork Reduction Act Officer, U.S. Customs and Border Protection, Office of Trade, Regulations and Rulings, Economic Impact Analysis Branch, 90 K Street NE, 10th Floor, Washington, DC 20229–1177.

FOR FURTHER INFORMATION CONTACT: Requests for additional PRA information should be directed to Seth Renkema, U.S. Customs and Border Protection, Office of Trade, Regulations and Rulings, Economic Impact Analysis Branch, 90 K Street NE, 10th Floor, Washington, DC 20229–1177, or via email CBP_PRA@cbp.dhs.gov. Please note that the contact information provided here is solely for questions regarding this notice. Individuals

seeking information about other CBP programs should contact the CBP National Customer Service Center at 877–227–5511, (TTY) 1–800–877–8339, or CBP website at www.cbp.gov/.

SUPPLEMENTARY INFORMATION: CBP invites the general public and other Federal agencies to comment on the proposed and/or continuing information collections pursuant to the Paperwork Reduction Act of 1995 (44 U.S.C. 3501 *et seq.*). This process is conducted in accordance with 5 CFR 1320.8. Written comments and suggestions from the public and affected agencies should address one or more of the following four points: (1) Whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information will have practical utility; (2) the accuracy of the agency’s estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used; (3) suggestions to enhance the quality, utility, and clarity of the information to be collected; and (4) suggestions to minimize the burden of the collection of information on those who are to respond, including through the use of appropriate automated, electronic, mechanical, or other technological collection techniques or other forms of information technology, *e.g.*, permitting electronic submission of responses. The comments that are submitted will be summarized and included in the request for approval. All comments will become a matter of public record.

Overview of This Information Collection

Title: e-Allegations Submission.

OMB Number: 1651–0131.

Form Number: None.

Current Actions: CBP proposes to extend the expiration date of this information collection. There is no change to the burden hours or to the information collected.

Type of Review: Extension (without change).

Affected Public: Businesses, Individuals.

Abstract: In the interest of detecting trade violations to customs laws, Customs and Border Protection (CBP) established the e-Allegations website to provide a means for concerned members of the trade community to confidentially report violations to CBP. The e-Allegations site allows the public to submit pertinent information that assists CBP in its decision whether or not to pursue the alleged violations by initiating an investigation. The information collected includes the

name, phone number and email address of the member of the trade community reporting the alleged violation. It also includes a description of the alleged violation, and the name and address of the potential violators. The e-Allegations website is accessible at <https://apps.cbp.gov/eallegations/>.

Estimated Number of Respondents: 1,600.

Estimated Number of Total Annual Responses: 1,600.

Estimated Time per Response: 15 minutes.

Estimated Total Annual Burden Hours: 400.

Dated: June 16, 2020.

Seth Renkema,

Branch Chief, Economic Impact Analysis Branch, U.S. Customs and Border Protection.

[FR Doc. 2020–13295 Filed 6–29–20; 8:45 am]

BILLING CODE 9111–14–P

DEPARTMENT OF THE INTERIOR

Bureau of Land Management

[LLES00000.L5110000.GF0000.LVEMM19M2070.19X]

Notice of Intent To Prepare an Environmental Impact Statement for the Twin Metals Project in the Superior National Forest, Lake and St. Louis Counties, Minnesota

AGENCY: Bureau of Land Management, Interior.

ACTION: Notice of intent.

SUMMARY: In compliance with the National Environmental Policy Act of 1969, as amended (NEPA), and the Federal Land Policy and Management Act of 1976 (FLPMA), as amended, the Bureau of Land Management (BLM) Northeastern States District, Milwaukee, Wisconsin, intends to prepare an Environmental Impact Statement (EIS) to analyze the potential impacts of issuing a proposed new preference right lease (MNES 57965) and approving a Mine Plan of Operation in the Superior National Forest in Lake and St. Louis Counties, Minnesota. The approval of a Mine Plan of Operation allows the lessee to access, and once other necessary permits are obtained, to mine federal minerals. The BLM will conduct a public scoping process, including public meetings. During this time, the public will be invited to submit comments.

DATES: The BLM will announce the dates of public scoping, including dates and locations of public meetings and the ways in which people may submit scoping comments, on its e-Planning