products, processes or services that are derived through SBIR funding: (2) determine if other measures of success defined within the NIH mission are being achieved; and (3) enhance NIH’s administration of the SBIR Program and the support that it provides to small business concerns. Overall, the NIH will use the evaluation results to assess the outcomes from NIH-supported SBIR awards. The evaluation results will provide OD with the information necessary to make quality improvements to the SBIR program and enhance program performance in generating significant outcomes. The Government Performance and Results Act of 1993 (GPRA) mandates that Federal programs improve their effectiveness and public accountability by focusing on results. The OMB developed the Program Assessment Rating Tool (PART) to monitor compliance with the GPRA and to rate federal programs for their effectiveness and ability to show results. It is anticipated that results from a second survey will assist NIH in demonstrating that it is meeting its GPRA goals for the NIH SBIR Program. Using an Internet survey OD will collect information Phase II SBIR awardees from fiscal years (FY) 2002 through 2006. The online survey will be implemented using Secure Socket Layer (SSL) encryption technology and password access. OD will use e-mail messages to advise awardees that they have been selected to participate in the survey.

Frequency of Response: One time.

Requests for Comments: Written comments and/or suggestions from the public and affected agencies are invited on one or more of the following points: (1) Whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information shall have practical utility; (2) The accuracy of the agency’s estimate of the burden (including hours and cost) of the proposed information collection; (3) Ways to enhance the quality, utility, and clarity of the information to be collected; and (4) Ways to minimize the burden of the collection of information on respondents, including through the use of appropriate automated, electronic, mechanical, or other technological collection techniques or other forms of information technology.

FOR FURTHER INFORMATION CONTACT: To request more information on the proposed project or to obtain a copy of the data collection plans and instruments, contact: Ms. Jo Anne Goodnight, NIH SBIR/STTR Program Coordinator, Rockledge I Bldg., Room 3538, 6705 Rockledge Drive, Bethesda, MD 20892–7910, or call non-toll-free number (301) 435–2688 or E-mail your request, including your address, to: jg128w@nih.gov.

Comments Due Date: Comments regarding this information collection are best assured of having their full effect if received on or before April 13, 2007.


Jo Anne Goodnight, Coordinator, Small Business Innovation Research/Small Business Technology Transfer Program Office of Extramural Programs, Office of the Director, National Institutes of Health.

[FR Doc. E7–2636 Filed 2–14–07; 8:45 am]

BILLING CODE 4140–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Substance Abuse and Mental Health Services Administration

Current List of Laboratories Which Meet Minimum Standards To Engage in Urine Drug Testing for Federal Agencies

AGENCY: Substance Abuse and Mental Health Services Administration, HHS.

ACTION: Notice.

SUMMARY: The Department of Health and Human Services (HHS) notifies Federal agencies of the laboratories currently certified to meet the standards of Subpart C of the Mandatory Guidelines for Federal Workplace Drug Testing Programs (Mandatory Guidelines). The Mandatory Guidelines were first published in the Federal Register on April 11, 1988 (53 FR 11970), and subsequently revised in the Federal Register on June 9, 1994 (59 FR 29908), on September 30, 1997 (62 FR 51118), and on April 13, 2004 (69 FR 19644).

A notice listing all currently certified laboratories is published in the Federal Register during the first week of each month. If any laboratory’s certification is suspended or revoked, the laboratory will be omitted from subsequent lists until such time as it is restored to full certification under the Mandatory Guidelines.

If any laboratory has withdrawn from the HHS National Laboratory Certification Program (NLCP) during the past month, it will be listed at the end, and will be omitted from the monthly listing thereafter.

This notice is also available on the Internet at http://workplace.samhsa.gov and http://www.drugfreeworkplace.gov.

FOR FURTHER INFORMATION CONTACT: Mrs. Giselle Hersh or Dr. Walter Vogl, Division of Workplace Programs, SAMHSA/CSAP, Room 2–1035, 1 Choke Cherry Road, Rockville, Maryland 20857; 240–276–2600 (voice), 240–276–2610 (fax).

SUPPLEMENTARY INFORMATION: The Mandatory Guidelines were developed in accordance with Executive Order 12564 and section 503 of Pub. L. 100–71. Subpart C of the Mandatory Guidelines, “Certification of Laboratories Engaged in Urine Drug Testing for Federal Agencies,” sets strict standards that laboratories must meet in order to conduct drug and specimen validity tests on urine specimens for Federal agencies. To become certified,

<table>
<thead>
<tr>
<th>Type of respondents</th>
<th>Estimated number of respondents</th>
<th>Estimated number of responses per respondent</th>
<th>Average burden hours per response</th>
<th>Estimated total annual burden hours requested</th>
</tr>
</thead>
<tbody>
<tr>
<td>For-profit small business concerns that have received an NIH SBIR Phase II award from (FY 2002–2006)</td>
<td>1000</td>
<td>1</td>
<td>0.5</td>
<td>500</td>
</tr>
</tbody>
</table>
an applicant laboratory must undergo three rounds of performance testing plus an on-site inspection. To maintain that certification, a laboratory must participate in a quarterly performance testing program plus undergo periodic, on-site inspections.

Laboratories which claim to be in the applicant stage of certification are not to be considered as meeting the minimum requirements described in the HHS Mandatory Guidelines. A laboratory must have its letter of certification from HHS/SAMHSA (formerly: HHS/NIDA) which attests that it has met minimum standards.

In accordance with Subpart C of the Mandatory Guidelines dated April 13, 2004 (69 FR 19644), the following laboratories meet the minimum standards to conduct drug and specimen testing.


Baptist Medical Center-Toxicology Laboratory, 9601 I–630, Exit 7, Little Rock, AR 72205–7299, 501–202–2783. (Formerly: Forensic Toxicology Laboratory Baptist Medical Center).

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


Doctors Laboratory, Inc., 2906 Julia Drive, Valdosta, GA 31602, 229–671–2281.

DrugScan, Inc., P.O. Box 2969, 1119 Mears Road, Warminster, PA 18974, 215–674–9310.


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, 10788 Roselle St., San Diego, CA 92121, 800–882–7272. (Formerly: Poisonlab, Inc.).

Laboratory Corporation of America Holdings, 550 17th Ave., Suite 300, Seattle, WA 98112, 206–923–7020 / 800–898–0180. (Formerly: DrugProof, Division of Dynacare/Laboratory of Pathology, LLC; Laboratory of Pathology of Seattle, Inc.; DrugProof, Division of Laboratory of Pathology of Seattle, Inc.).

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

Marshfield Laboratory, Forensic Toxicology Laboratory, 1000 North Oak Ave., Marshfield, WI 54449, 715–389–3734/800–331–3734. (Formerly: MAXXAM Analytics Inc., * 6740 Campbell Road, Mississauga, ON, Canada L5N 2L8, 905–817–5700. (Formerly: NOVAMANN (Ontario), Inc.).


Meritor Laboratories, 36 South Brooks St., Madison, WI 53715, 608–267–6225. (Formerly: General Medical Laboratories).

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

Minneapolis Veterans Affairs Medical Center, Forensic Toxicology Laboratory, 1 Veterans Drive, Minneapolis, MN 55417, 612–725–2088.


One Source Toxicology Laboratory, Inc., 1213 Genoa-Red Bluff, Pasadena, TX 77504, 888–747–3774. (Formerly: University of Texas Medical Branch, Clinical Chemistry Division; UTMB Pathology-Toxicology Laboratory).


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


Quest Diagnostics Incorporated, 3175 Presidential Dr., Atlanta, GA 30340, 770–452–1590/800–729–6432. (Formerly: SmithKline Beecham Clinical Laboratories; SmithKline Biotechnology Laboratories).


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


South Bend Medical Foundation, Inc., 530 N. Lafayette Blvd., South Bend, IN 46601, 574–234–4176 x276.

Southwest Laboratories, 4645 E. Cotton Center Boulevard, Suite 177, Phoenix,
DEPARTMENT OF HOMELAND SECURITY

Bureau of Customs and Border Protection

Proposal Collection; Comment Request; Distribution of Continued Dumping and Subsidy Offset

ACTION: Notice and request for comments.

SUMMARY: As part of its continuing effort to reduce paperwork and respondent burden, the Bureau of Customs and Border Protection (CBP) invites the general public and other Federal agencies to comment on an information collection requirement concerning the Distribution of Continued Dumping and Subsidy Offset to Affected Domestic Procedures. This request for comment is being made pursuant to the Paperwork Reduction Act of 1995 (Pub. L. 104–13; 44 U.S.C. 3505(c)(2)).

DATES: Written comments should be received on or before April 16, 2007, to be assured of consideration.

ADDRESSES: Direct all written comments to Bureau of Customs and Border Protection, Information Services Group, Room 3.2.C., 1300 Pennsylvania Ave., NW., Washington, DC 20229.

FOR FURTHER INFORMATION CONTACT: Requests for additional information should be directed to Bureau of Customs and Border Protection, Attn.: Tracey Denning, Room 3.2.C., 1300 Pennsylvania Avenue, NW., Washington, DC 20229, Tel. (202) 344–1429.

SUPPLEMENTARY INFORMATION: CBP invites the general public and other Federal agencies to comment on proposed and/or continuing information collections pursuant to the Paperwork Reduction Act of 1995 (Pub. L. 104–13; 44 U.S.C. 3505(c)(2)). The comments should address: (a) Whether the collection of information is necessary for the proper performance of the functions of the agency, including whether the information shall have practical utility; (b) the accuracy of the agency’s estimates of the burden of the collection of information; (c) ways to enhance the quality, utility, and clarity of the information to be collected; (d) ways to minimize the burden including the use of automated collection techniques or the use of other forms of information technology; and (e) estimates of capital or start-up costs and costs of operations, maintenance, and purchase of services to provide information. The comments that are submitted will be summarized and included in the CBP request for Office of Management and Budget (OMB) approval. All comments will become a matter of public record. In this document, CBP is soliciting comments concerning the following information collection:

Title: Distribution of Continued Dumping and Subsidy Offset to Affected Domestic Producers.

OMB Number: 1651–0086.

Form Number: N/A.

Abstract: The collection of information is required to implement the duty preference provisions of the Continued Dumping and Subsidy Offset Act of 2000, by prescribing the administrative procedures under which anti-dumping and countervailing duties are assessed on imported products.

Current Actions: This submission is being submitted to extend the expiration date.

Type of Review: Extension (without change).

Affected Public: Businesses.

Estimated Number of Respondents: 2000.

Estimated Time per Respondent: 1 hour.

Estimated Total Annual Burden Hours: 2000.

Estimated Total Annualized Cost on the Public: N/A.


Tracey Denning,

Agency Clearance Officer, Information Services Group.

[FR Doc. E7–2655 Filed 2–14–07; 8:45 am]

BILLING CODE 9111–14–P

DEPARTMENT OF HOMELAND SECURITY

Bureau of Customs and Border Protection

Proposed Collection; Comment Request; Accreditation of Commercial Testing Laboratories; Approval of Commercial Gaugers

ACTION: Notice and request for comments.

SUMMARY: As part of its continuing effort to reduce paperwork and respondent burden, CBP invites the general public and other Federal agencies to comment on an information collection requirement concerning the Accreditation of Commercial Testing Laboratories and Approval of Commercial Gaugers. This request for comment is being made pursuant to the Paperwork Reduction Act of 1995 (Pub. L. 104–13; 44 U.S.C. 3505(c)(2)).