internally and for sale to other companies.

Any other such applicant and any person who is presently registered with DEA to manufacture such substances may file comments or objections to the issuance of the proposed registration.

Any such comments or objections may be addressed, in quintuplicate, to the Deputy Assistant Administrator, Office of Diversion Control, Drug Enforcement Administration, United States Department of Justice, Washington, DC 20537. Attention: DEA Federal Register Representative (CCR), and must be filed no later than April 17, 2002.


Laura M. Nagel,
Deputy Assistant Administrator, Office of Diversion Control, Drug Enforcement Administration.

[FR Doc. 02–6324 Filed 3–15–02; 8:46 am]
BILLING CODE 4410–09–M

DEPARTMENT OF JUSTICE
Drug Enforcement Administration

Importation of Controlled Substances; Notice of Application

Pursuant to section 1006 of the Controlled Substances Import and Export Act (21 U.S.C. 958(i)), the Attorney General shall, prior to issuing a registration under this Section to a bulk manufacturer of a controlled substance in Schedule I or II and prior to issuing a registration under section 1002(a) authorizing the importation of such a substance, provide manufacturers holding registrations for the bulk manufacture of the substance an opportunity for a hearing.

Therefore, in accordance with 1301.34 of Title 21, Code of Federal Regulations (CFR), notice is hereby given that on September 5, 2001, Roxane Laboratories, Inc., 1809 Wilson Road, PO. Box 16532, Columbus, Ohio 43216–6532, made application by renewal to the Drug Enforcement Administration to be registered as an importer of cocaine (9041), a basic class of controlled substance listed in Schedule II.

The firm plans to import cocaine to the United States Department of Justice, Office of Diversion Control, Drug Enforcement Administration to be manufactured for sale to other companies.

Any other such applicant and any person who is presently registered with DEA to manufacture such substances may file comments or objections to the issuance of the proposed registration.

Any such comments or objections may be addressed, in quintuplicate, to the Deputy Assistant Administrator, Office of Diversion Control, Drug Enforcement Administration, United States Department of Justice, Washington, DC 20537. Attention: DEA Federal Register Representative (CCR), and must be filed no later than April 17, 2002.

This procedure is to be conducted simultaneously with and independent of the procedures described in 21 CFR 1301.34(b), (c), (d), (e), and (f). As noted in a previous notice at 40 FR 43745–46 (September 23, 1975), all applicants for registration to import the basic class of any controlled substance in Schedule I or II are and will continue to be required to demonstrate to the Deputy Assistant Administrator, Office of Diversion Control, Drug Enforcement Administration that the requirements for such registration pursuant to 21 U.S.C. 958(a), 21 U.S.C. 823(a), and 21 CFR 1301.34(a), (b), (c), (d), (e), and (f) are satisfied.

Dated: March 7, 2002.

Laura M. Nagel,
Deputy Assistant Administrator, Office of Diversion Control, Drug Enforcement Administration.

[FR Doc. 02–6321 Filed 3–15–02; 8:45 am]
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DEPARTMENT OF LABOR

Occupational Safety and Health Administration

[Docket No. ICR–1218–0072(2002)]

Hazard Communication Standard; Extension of the Office of Management and Budget’s (OMB) Approval of Information-Collection (Paperwork) Requirements

AGENCY: Occupational Safety and Health Administration (OSHA), Labor.

ACTION: Request for comments.

SUMMARY: OSHA solicits comments concerning its proposal to decrease the total burden hour estimates for, and to extend OMB approval of, the collection-of-information requirements specified by the Hazard Communication Standard (HCS) (29 CFR 1910.1200; 1915.1200; 1917.28; 1918.90; 1926.59; 1928.21).1

1Based on its assessment of the paperwork requirements contained in this standard, the Agency estimates that the total burden hours decreased compared to its previous burden-hour estimate. Under this notice, OSHA is not proposing to revise these paperwork requirements in any manner, only to decrease the burden-hour estimate imposed by the existing paperwork requirements.
Communication Standard ensures that the hazards of all chemicals produced or imported are evaluated and that information concerning their hazards is transmitted to employees and downstream employers. The Standard requires chemical manufacturers and importers to evaluate chemicals they produce or import to determine if they are hazardous; for those chemicals determined to be hazardous, material safety data sheets and warning labels must be developed. Employers are required to establish hazard communication programs, to transmit information on the hazards of chemicals to their employees by means of labels on containers, material safety data sheets and training programs.

Implementation of these collection of information requirements will ensure all employees have the “right-to-know” the hazards and identities of the chemicals they work with and will reduce the incidence of chemically-related occupational illnesses and injuries.

II. Special Issues for Comment

OSHA has particularly interest in comments on the following issues:

- Whether the information collection requirements are necessary for the proper performance of the Agency’s functions, including whether the information is useful;
- The accuracy of OSHA’s estimate of the burden (time and costs) of the information collection requirements, including the validity of the methodology and assumptions used;
- The quality, utility, and clarity of the information collected; and
- Ways to minimize the burden on employers who must comply, for example, by using automated, or other technological information-collection and -transmission techniques.

III. Proposed Actions

OSHA is requesting to decrease the existing burden-hour estimate for, and to extend OMB approval of, the collection-of-information requirements specified by the Standard. In this regard, the Agency is requesting to decrease the current burden hour estimate from 7,560,232 hours to 7,498,766 hours, a total decrease of 61,466 hours. Based on more recent data, the Agency reduced the number of establishments, shipped containers, and in-plant containers used to calculate the burden hour and cost estimates. OSHA will summarize the comments submitted in response to this notice, and will include this summary in its request to OMB to extend its approval of these information-collection requirements.

IV. Authority and Signature

John L. Henshaw, Assistant Secretary of Labor for Occupational Safety and Health, directed the preparation of this notice. The authority for this notice is the Paperwork Reduction Act of 1995 (44 U.S.C. 3506), and Secretary of Labor’s Order No. 3-2000 (65 FR 50017).

Signed at Washington, DC on March 8, 2002.
John L. Henshaw, Assistant Secretary of Labor.
[FR Doc. 02–6449 Filed 3–15–02; 8:45 am]
BILLING CODE 4510–26–M

DEPARTMENT OF LABOR

Occupational Safety and Health Administration
[Docket No. NRTL3–92]

TUV Rheinland of North America, Inc., Renewal of Recognition

AGENCY: Occupational Safety and Health Administration (OSHA), Labor.

ACTION: Notice.

SUMMARY: This notice announces the Agency’s final decision on the application of the TUV Rheinland of North America, Inc. (TUV), for renewal of its recognition as a Nationally Recognized Testing Laboratory under 29 CFR 1910.7.

EFFECTIVE DATE: The renewal becomes effective on March 18, 2002 and will be valid until March 19, 2007, unless terminated or modified prior to that date, in accordance with 29 CFR 1910.7.

FOR FURTHER INFORMATION CONTACT: Bernard Pasquet or Sherrey Nicolas, Office of Technical Programs and Coordination Activities, NRTL Program, Occupational Safety and Health Administration, U.S. Department of Labor, 200 Constitution Avenue, NW., Room N3653, Washington, DC 20210, or phone (202) 693–2110.

SUPPLEMENTARY INFORMATION:

Notice of Final Application

The Occupational Safety and Health Administration (OSHA) hereby gives notice of the renewal of recognition of TUV Rheinland of North America, Inc. (TUV), as a Nationally Recognized Testing Laboratory (NRTL). The NRTL’s scope of recognition may be found in OSHA’s informational web page for the NRTL (http://www.osha-slc.gov/dts/otpca/nrtl/tuv.html). OSHA recognition of an NRTL signifies that the organization has met the legal requirements in Section 1910.7 of Title 29, Code of Federal Regulations (29 CFR 1910.7). Recognition is an acknowledgment that the organization can perform independent safety testing and certification of the specific products covered within its scope of recognition and is not a delegation or grant of government authority. As a result of recognition, employers may use products “properly certified” by the NRTL to meet OSHA standards that require testing and certification.

The Agency processes applications by an NRTL for initial recognition or for expansion or renewal of this recognition following requirements in Appendix A to 29 CFR 1910.7. This appendix requires that the Agency publish two notices in the Federal Register in processing an application. In the first notice, OSHA announces the application and provides its preliminary finding and, in the second notice, the Agency provides its final decision on the application. These notices set forth the NRTL’s scope of recognition or modifications of that scope.

TUV Rheinland of North America, Inc., is a privately held Product Safety and Quality Assurance Testing firm with offices throughout the United States and Canada. TUV is wholly owned by TUV Rheinland e. V. of Cologne, Germany. TUV is a U.S. corporation incorporated in the state of Delaware in 1983.

TUV received its recognition as an NRTL on August 16, 1995 (60 FR 42594), for a period of five years ending August 16, 2000. Appendix A to 29 CFR 1910.7 stipulates that the period of recognition of an NRTL is five years and that an NRTL may renew its recognition by applying not less than nine months, nor more than one year before the expiration date of its current recognition. TUV submitted its renewal...