tolerances will utilize 2.2% of the RfD for the U.S. population.

G. Effects on the Immune and Endocrine Systems

Lifespan, and multigenerational studies on mammals, and acute and subchronic studies on aquatic organisms and wildlife did not reveal any definite immune or endocrine effects. An immunotoxicity study in rats at 0, 1.25, 5 and 15 mg/kg/day with a NOAEL of 5 mg/kg/day based on decreased primary antibody (igM) response to sheep red blood cells; decreased absolute and relative thymus weights; decreased body weight, food consumption and food efficiency at the high dose level. The LOEL is 15 mg/kg/ day.

Any endocrine related effects would have been detected in this definitive array of required tests. The probability of any such effect due to agricultural uses of AVG is considered negligible.

H. Existing Tolerances

Tolerances have been established for the residues of AVG in or on the following food commodities:

Commodity	Parts per million
Apples	0.08
Fruit, stone, group 12, (except cherry)	0.170
Pears	0.08

I. International Tolerances

There are no Codex maximum residue limits for use of aviglycine HCl on apples or pears, or on any other crop.

[FR Doc. 05–8791 Filed 5–3–05; 8:45 am] BILLING CODE 6560–50–S

ENVIRONMENTAL PROTECTION AGENCY

[OPPT-2005-0026; FRL-7713-6]

Approval of Test Marketing Exemption for a Certain New Chemical

AGENCY: Environmental Protection Agency (EPA). **ACTION:** Notice.

ACTION: Notice.

SUMMARY: This notice announces EPA's approval of an application for test marketing exemption (TME) under section 5(h)(1) of the Toxic Substances Control Act (TSCA) and 40 CFR 720.38. EPA has designated this application as TME–05–3. The test marketing conditions are described in the TME application and in this notice.

DATES: Approval of this TME is effective April 27, 2005.

FOR FURTHER INFORMATION CONTACT: For general information contact: Colby Lintner, Regulatory Coordinator, Environmental Assistance Division (7408M), Office of Pollution Prevention and Toxics, Environmental Protection Agency, 1200 Pennsylvania Ave., NW., Washington, DC 20460–0001; telephone number: (202) 554–1404; e-mail address: TSCA-Hotline@epa.gov.

For technical information contact: Adella Underdown, Program Manager, Chemical Control Division (7405M), Office of Pollution Prevention and Toxics, Environmental Protection Agency, 1200 Pennsylvania Ave., NW., Washington, DC 20460–0001; telephone number: (202) 564–9364; e-mail address: underdown.adella@epa.gov. SUPPLEMENTARY INFORMATION:

I. General Information

A. Does this Action Apply to Me?

This action is directed in particular to the chemical manufacturer and/or importer who submitted the TME to EPA. This action may, however, be of interest to the public in general. Since other entities may also be interested, the Agency has not attempted to describe all the specific entities that may be affected by this action. If you have any questions regarding the applicability of this action to a particular entity, consult the technical person listed under FOR FURTHER INFORMATION CONTACT.

B. How Can I Get Copies of this Document and Other Related Information?

1. Docket. EPA has established an official public docket for this action under docket identification (ID) number OPPT-2005-0026. The official public docket consists of the documents specifically referenced in this action, any public comments received, and other information related to this action. Although a part of the official docket, the public docket does not include Confidential Business Information (CBI) or other information whose disclosure is restricted by statute. The official public docket is the collection of materials that is available for public viewing at the EPA Docket Center, Rm. B102-Reading Room, EPA West, 1301 Constitution Ave., NW., Washington, DC. The EPA Docket Center is open from 8:30 a.m. to 4:30 p.m., Monday through Friday, excluding legal holidays. The EPA Docket Center Reading Room telephone number is (202) 566-1744 and the telephone number for the OPPT Docket. which is located in EPA Docket Center, is (202) 566-0280.

2. *Electronic access.* You may access this **Federal Register** document electronically through the EPA Internet under the "**Federal Register**" listings at *http://www.epa.gov/fedrgstr/.*

An electronic version of the public docket is available through EPA's electronic public docket and comment system, EPA Dockets. You may use EPA Dockets at http://www.epa.gov/edocket/ to submit or view public comments, access the index listing of the contents of the official public docket, and to access those documents in the public docket that are available electronically. Although not all docket materials may be available electronically, you may still access any of the publicly available docket materials through the docket facility identified in Unit I.B.1. Once in the system, select "search," then key in the appropriate docket ID number.

II. What is the Agency's Authority for Taking this Action?

Section 5(h)(1) of TSCA and 40 CFR 720.38 authorizes EPA to exempt persons from premanufacture notification (PMN) requirements and permit them to manufacture or import new chemical substances for test marketing purposes, if the Agency finds that the manufacture, processing, distribution in commerce, use, and disposal of the substances for test marketing purposes will not present an unreasonable risk of injury to health or the environment. EPA may impose restrictions on test marketing activities and may modify or revoke a test marketing exemption upon receipt of new information which casts significant doubt on its finding that the test marketing activity will not present an unreasonable risk of injury.

III. What Action is the Agency Taking?

EPA approves the above-referenced TME. EPA has determined that test

marketing the new chemical substance, under the conditions set out in the TME application and in this notice, will not present an unreasonable risk of injury to health or the environment.

IV. What Restrictions Apply to this TME?

The test market time period, production volume, number of customers, and use must not exceed specifications in the application and this notice. All other conditions and restrictions described in the application and in this notice must also be met.

TME-05-03.

Date of Receipt: March 14, 2005. Notice of Receipt: April 8, 2005 (70 FR 18013) (FRL–7708–8).

Applicant: CBI. Chemical: (G) Soy Polyol. Use: (G) Polyurethane's market Production Volume: CBI. Number of Customers: CBI. Test Marketing Period: CBI.

The following additional restrictions apply to this TME. A bill of lading accompanying each shipment must state that the use of the substance is restricted to that approved in the TME. In addition, the applicant shall maintain the following records until 5 years after the date they are created, and shall make them available for inspection or copying in accordance with section 11 of TSCA:

1. Records of the quantity of the TME substance produced and the date of manufacture.

2. Records of dates of the shipments to each customer and the quantities supplied in each shipment.

3. Copies of the bill of lading that accompanies each shipment of the TME substance.

V. What was EPA's Risk Assessment for this TME?

EPA identified no significant health or environmental concerns for the test market substance. Therefore, the test market activities will not present an unreasonable risk of injury to human health or the environment.

VI. Can EPA Change Its Decision on this TME in the Future?

Yes. The Agency reserves the right to rescind approval or modify the conditions and restrictions of an exemption should any new information that comes to its attention cast significant doubt on its finding that the test marketing activities will not present an unreasonable risk of injury to human health or the environment.

List of Subjects

Environmental protection, Test marketing exemptions.

Dated: April 27, 2005.

Anna Coutlakis,

Acting Chief, New Chemicals Prenotice Management Branch, Office of Pollution Prevention and Toxics.

[FR Doc. 05–8790 Filed 5–3–05 8:45 am] BILLING CODE 6560–50–S

FEDERAL COMMUNICATIONS COMMISSION

Public Information Collections Approved by Office of Management and Budget

April 26, 2005.

SUMMARY: The Federal Communications Commission (FCC) has received Office of Management and Budget (OMB) approval for the following public information collections pursuant to the Paperwork Reduction Act of 1995, Public Law 104–13. An agency may not conduct or sponsor and a person is not required to respond to a collection of information unless it displays a currently valid control number.

FOR FURTHER INFORMATION CONTACT: Paul J. Laurenzano, Federal Communications Commission, 445 12th Street, SW., Washington DC 20554, (202) 418–1359 or via the Internet at *plaurenz@fcc.gov.* SUPPLEMENTARY INFORMATION:

OMB Control No.: 3060–1081. OMB Approval Date: 04/15/2005. Expiration Date: 04/30/2008. Title: Federal-State Joint Board on Universal Service Petitions for

Designations as Eligible

Telecommunications Carriers, CC

Docket No. 96–45.

Form No.: N/A.

Estimated Annual Burden: 22 responses; 176 total annual burden hours; approximately 8 hours average per respondent.

Needs and Uses: In the Virginia Cellular Order (FCC 03-338), the Commission stated as part future Eligible Telecommunications Carriers (ETC) designation orders, each designated ETC will be required to submit records and documentation on an annual basis. In particular, ETCs will be required to report: (1) Progress towards meeting infrastructure buildout plans; (2) the number of consumer complaints per 1,000 handsets; and (3) information detailing the number of unfulfilled requests for service from potential customers for a twelve month period. This information collection is necessary to ensure that each ETC satisfies its obligation under section 214(e) of the Communications Act of 1934, as amended, to provide services supported by the universal service

mechanism throughout the areas for which each ETC is designated.

OMB Control No.: 3060–0814. OMB Approval date: 03/16/2005. Expiration Date: 03/31/2008. Title: Section 54.301, Local Switching

Support and Local Switching Support Data Collection Form and Instructions.

Form No.: N/A.

Estimated Annual Burden: 160 responses; 3,324 total annual burden hours; .5–24 hours average response time per respondent.

Needs and Uses: Pursuant to section 54.301, each incumbent local exchange carrier that is not a member of the NECA common line tariff, that has been designated an eligible telecommunications carriers, and that serves a study area with 50,000 or fewer access lines shall, for each study area, provide the Administrator with the projected total unseparated dollar amount assigned to each account in section 54.301(b). Average schedule companies are required to file information pursuant to section 54.301(f). Both respondents must provide true-up data. The data is necessary to calculate certain revenue requirement.

OMB Control No.: 3060–0512. OMB Approval date: 4/15/2005. Expiration Date: 4/30/2008. Title: The ARMIS Annual Summary Report.

Form No.: FCC 43–01.

Estimated Annual Burden: 124 responses; 11,036 total annual burden hours; 89 hours per respondent.

Needs and Uses: The Annual Summary Report contains financial and operating data and is used to monitor the incumbent local exchange carrier industry and to perform routine analyses of costs and revenues on behalf of the Commission.

OMB Control No.: 3060–0511. OMB Approval date: 4/15/2005. Expiration Date: 4/30/2008. Title: ARMIS Access Report. Form No.: FCC Report 43–04. Estimated Annual Burden: 82

responses; 12,546 total annual burden hours; 153 hours average per respondent.

Needs and Uses: The Access Report is needed to administer the Commission's accounting, jurisdicational separations and access charge rule; to analyze revenue requirements and rates of return, and to collect financial data from Tier 1 incumbent local exchange carriers.

OMB Control No.: 3060–0470. *OMB Approval date:* 3/25/2005. *Expiration Date:* 3/31/2008.