available for public inspection. This filing may also be viewed on the web at http://www.ferc.gov using the "RIMS" link, select "Dockett" and follow the instructions (call 202–208–2222 for assistance). Comments, protests and interventions may be filed electronically via the Internet in lieu of paper. See, 18 CFR 385.2001(a)(1)(iii) and the instructions on the Commission's web site under the "e-Filing" link.

Linwood A. Watson, Jr.,

Acting Secretary.
[FR Doc. 02–466 Filed 1–8–02; 8:45 am]
BILLING CODE 6717–01–P

DEPARTMENT OF ENERGY

Federal Energy Regulatory Commission

[Docket No. CP02-025-000]

Copiah County Storage Company; Notice of Route and Site Review

January 3, 2002.

On January 16, 2002, the staff of the Office of Energy Projects (OEP) will conduct a site review of the proposed Copiah Storage Project. The Copiah Storage Project facilities are proposed for construction by Copiah County Storage Company (Copiah). The proposed compressor site and natural gas storage cavern site, located in Copiah County, Mississippi, will be reviewed on January 16, 2002. Representatives of Copiah will accompany the OEP staff.

Anyone interested in attending the route and site review or obtaining further information may contact the Commission's Office of External Affairs at (202) 208–1088. Attendees must provide their own transportation.

Linwood A. Watson, Jr.,

Acting Secretary.

[FR Doc. 02–463 Filed 1–8–02; 8:45 am] BILLING CODE 6717–01–P

DEPARTMENT OF ENERGY

Federal Energy Regulatory Commission

[Docket No. ER02-517-000]

UtiliGroup, Inc.; Notice of Filing

January 3, 2002.

Take notice that on December 10, 2001, UtiliGroup, Inc. (UtiliGroup), tendered for filing with the Federal Energy Regulatory Commission (Commission) a Petition For Acceptance of Initial Rate Schedule, Waivers and Blanket Authority. The Petition request

acceptance of UtiliGroup Rate Schedule FERC No. 1, under which UtiliGroup will engage in wholesale electric power and energy transactions as a marketer.

Any person desiring to be heard or to protest such filing should file a motion to intervene or protest with the Federal Energy Regulatory Commission, 888 First Street, N.E., Washington, DC 20426, in accordance with Rules 211 and 214 of the Commission's Rules of Practice and Procedure (18 CFR 385.211 and 385.214). All such motions and protests should be filed on or before the comment date. Protests will be considered by the Commission to determine the appropriate action to be taken, but will not serve to make protestants parties to the proceedings. Any person wishing to become a party must file a motion to intervene. Copies of this filing are on file with the Commission and are available for public inspection. This filing may also be viewed on the Commission's web site at http://www.ferc.gov using the "RIMS" link, select "Docket#" and follow the instructions (call 202-208-2222 for assistance). Comments, protests and interventions may be filed electronically via the internet in lieu of paper. See, 18 CFR 385.2001(a)(1)(iii) and the instructions on the Commission's web site under the "e-filing" link.

Comment Date: January 11, 2002.

Linwood A. Watson, Jr.,

Acting Secretary.

[FR Doc. 02–464 Filed 1–8–02; 8:45 am] BILLING CODE 6717–01–P

ENVIRONMENTAL PROTECTION AGENCY

[FRL-7127-2]

Integrated Risk Information System (IRIS); Announcement of 2002 Program; Request for Information

AGENCY: Environmental Protection Agency.

ACTION: Notice; announcement of IRIS 2002 program and request for scientific information on health effects that may result from exposure to chemical substances.

SUMMARY: IRIS is an Environmental Protection Agency (EPA) data base that contains EPA scientific consensus positions on human health effects that may result from chronic exposure to chemical substances in the environment. On February 22, 2001, EPA announced the 2001 IRIS agenda and solicited scientific information from the public for consideration in assessing health effects from specific chemical

substances (66 FR 11165). Most of the health assessments listed in the notice are in progress or near completion. Today, EPA is adding some additional health assessments to the IRIS agenda. This notice describes the Agency's plans, and solicits scientific data and evaluations for consideration in EPA's new assessments. Additional new assessments may be announced in the **Federal Register** later this year.

DATES: Please submit any response to this notice in the form of an initial "submission inventory" in accordance with the instructions in this notice by March 11, 2002.

ADDRESSES: A "submission inventory" should be sent to the IRIS Submission Desk in accordance with the instructions provided under "Submission of Information" in this notice.

FOR FURTHER INFORMATION: For information on the IRIS progra

information on the IRIS program, contact Amy Mills, National Center for Environmental Assessment, (mail code 8601D), U.S. Environmental Protection Agency, Washington, DC 20460, or call (202) 564–3204, or send electronic mail inquiries to mills.amy@epa.gov. For general questions about access to IRIS, or the content of IRIS, please call the IRIS Hotline at (301) 345–2870 or send electronic mail inquiries to hotline.iris@epa.gov.

SUPPLEMENTARY INFORMATION:

Background

IRIS is an EPA data base containing Agency consensus scientific positions on potential adverse human health effects that may result from chronic (or lifetime) exposure to chemical substances found in the environment. IRIS currently provides health effects information on over 500 specific chemical substances.

IRIS contains chemical-specific summaries of qualitative and quantitative health information in support of the first two steps of the risk assessment process, i.e., hazard identification and dose-response evaluation. IRIS information includes the reference dose for noncancer health effects resulting from oral exposure, the reference concentration for non-cancer health effects resulting from inhalation exposure, and the carcinogen assessment for both oral and inhalation exposure. Combined with specific situational exposure assessment information, the summary health hazard information in IRIS may be used as a source in evaluating potential public health risks from environmental contaminants.

The IRIS Program

EPA's process for developing IRIS consists of: (1) An annual Federal Register announcement of EPA's IRIS agenda and call for scientific information from the public on the selected chemical substances, (2) a search of the current literature, (3) development of health assessments and draft IRIS summaries, (4) peer review within EPA, (5) peer review outside EPA, (6) EPA consensus review and management approval, (7) preparation of final IRIS summaries and supporting documents, and (8) entry of summaries and supporting documents into the IRIS data base.

This notice provides: (1) A list of the IRIS assessments completed in FY 2001 and early FY 2002, (2) a list of the IRIS assessments in progress that the Agency expects to complete in FY 2002 or FY 2003, (3) a list of the IRIS assessments in progress that the Agency expects to complete in FY 2004 or FY 2005, (4) a list of the IRIS assessments announced in the previous IRIS agenda (66 FR 11165) that have been discontinued, (5) information about EPA's IRIS "needs assessment" report underway, (6) a list of the new assessments beginning in FY 2002, and (7) instructions to the public for submitting scientific information to EPA pertinent to the development of IRIS assessments.

Assessments Completed in FY 2001 and Early FY 2002

The following assessments were completed and entered into IRIS in FY 2001 and early FY 2002. These assessments were listed in the **Federal Register** of February 22, 2001. All health endpoints, cancer and noncancer, were assessed unless otherwise noted. Where information was available, both qualitative and quantitative assessments were developed.

Name	CAS No.
Bromate	7758–01–2 10049–04–4 7758–19–2 67–66–3 77–47–4
endpoints)	22967–92–6 74–87–3 91–22–5

Assessments in Progress—Completion Planned for FY 2002 or FY 2003

The following assessments are underway or generally complete, and are planned for entry into IRIS in FY 2002 or FY 2003. These assessments were announced in the February 22, 2001, Federal Register. All health

endpoints, cancer and noncancer, are being assessed unless otherwise noted. For all endpoints assessed, both qualitative and quantitative assessments are being developed where information is available. Pesticides denoted with an asterisk (*) will have only oral reference dose and carcinogenicity endpoints assessed.

Name	CAS No.
Acetaldehyde	75-07-0
Acetone	67–64–1
Acrolein	107-02-8
Alachlor*	15972-60-8
Ammonium perchlorate (and	
other perchlorate salts)	7790–98–9
Antimony and compounds	7440-36-0
Azinphos Methyl*	86-50-0
Benzene (noncancer endpoints)	71-43-2
Benzo(a)pyrene	50-32-8
Bromoxvnil*	1689-84-5
Boron	7440–42–8
1,3-Butadiene	106–99–0
Cadmium	7440–43–9
Carbon tetrachloride	56–23–5
Chloroethane	75–00–3
Chloroform (inhalation route)	67–66–3
Chloroprene	126-99-8
Chlorothalonil*	1897–45–6
Chlorpyrifos*	2921–88–2
Copper	7440–50–8
Cyclohexane	110-82-7
Diazinon*	333-41-5
Dichloroacetic acid	79–43–6
1,2-Dichlorobenzene	95–50–1 541–73–1
1,3-Dichlorobenzene	106-46-7
1,4-Dichlorobenzene 1,1-Dichloroethylene	75–35–4
Diesel exhaust	75–35–4 [N.A.]
Di(2-ethylhexyl)phthalate	117–81–7
Diflubenzuron	35367–38–5
Ethanol	64–17–5
Ethion*	563-12-2
Ethylbenzene	100-41-4
Ethylene dibromide	106–93–4
Ethylene dichloride	107-06-2
Ethylene oxide	75–21–8
Formaldehyde	50-00-0
Glyphosate*	1071-83-6
Hexachlorobutadiene	87-68-3
Hydrogen sulfide	7783-06-4
Isopropanol Metolachlor*	67–63–0
Metolachlor*	51218-45-2
Methidathion*	950–37–8
Methyl isobutyl ketone (MIBK)	108–10–1
Methyl parathion*	298–00–0
Methyl tert-butyl ether (MTBE)	1634–04–4
Mirex	2385–85–5
Nickel (soluble salts)	[N.A.]
Nitrobenzene	98–95–3
Pendimethalin*	40487-42-1
Phenol	108-95-2
Pebulate*	1114–71–2
Pentachlorophenol	87–86–5
Phosgene	75–44–5
Polychlorinated biphenyls	1226 26 2
(PCBs-noncancer endpoints)	1336–36–3
Refractory ceramic fibers	[N.A.] 100–42–5
Styrene	1746-01-6
Tetrachloroethylene	1740-01-6
(perchloroethylene)	127–18–4
Tetrahydrofuran	109-99-9
rottattyatotatatt	103-33-3

Name	CAS No.
Toluene	108-88-3 2303-17-5 55335-06-3 79-01-6 7440-61-1 108-05-4 1330-20-7 7440-66-6

Assessments in Progress—Completion Planned for FY 2004 or FY 2005

The following assessments in progress have been delayed and are now expected for completion in FY 2004 or FY 2005:

Name	CAS No.
Acrylamide	79–06–1 1332–21–4
(RDX)	121–82–4 67–56–1 14808–60–7

IRIS summaries and support documents for all substances listed above will be provided on the IRIS web site at www.epa.gov/iris as they are completed. This publicly available web site is EPA's primary location for IRIS documents. In addition, external peer review drafts of IRIS documents can be found during their peer review periods via the "What's New" page of the IRIS web site. Interested parties should check the "What's New" page frequently for the availability of these drafts.

Assessments Discontinued

The following assessments have been removed from the IRIS agenda for FY 2002, but may be reconsidered at a later date:

Name	CAS No.
Arsenic, inorganic Bisphenol-A Hexachlorobenzene Methylene chloride Toxaphene	7440-38-2 80-05-7 118-74-1 75-09-2 8001-35-2

IRIS "Needs Assessment"

On July 20, 2001, EPA published a **Federal Register** notice (66 FR 37958) requesting public input to compile a "needs assessment" for planning the IRIS program. This notice requested that the public identify those chemical substances for which assessments either need to be added to IRIS or updated. The responses were considered along with EPA program priorities in the development of new starts for the FY 2002 agenda below. The notice also requested input on whether other types of evaluations are needed on IRIS, such

as toxicological evaluations for health effects associated with less-than-lifetime (i.e., acute or subchronic) exposure durations. The notice also requested input on what priority any new type of evaluation should have compared to evaluation of health effects associated with chronic exposures.

Further, the notice asked whether or how EPA should work with external parties, such as other government agencies, industries, or other organizations to develop health assessments that may be used as supporting documents for IRIS. A pilot effort to provide dialogue and feedback to external parties developing health assessments for IRIS was described in the February 22, 2001 notice, and preceding IRIS Federal Register notices. Of the six pilot efforts discussed, four are still in progress and two are discontinued. In FY 2002, EPA will continue to evaluate its experience with the current efforts to determine process efficiency and quality of the documents produced.

A separate "IRIS Needs Assessment" report will be made available on the IRIS web site when it is completed.

Information Requested on New Assessments for FY 2002

EPA will continue building and updating the IRIS data base. The Agency recognizes that a number of the assessments on IRIS need updating to incorporate new scientific information and methodologies. Further, many additional substances are candidates for adding to IRIS. However, due to limited resources in the Agency to address the spectrum of needs, EPA developed a list of priority substances for attention beginning in FY 2002. The following substances are priorities for IRIS due to one or more reasons: (1) Agency statutory, regulatory, or program implementation needs; (2) new scientific information or methodology is available that might significantly change current IRIS information; (3) interest to other levels of government or the public, including interest expressed via responses to 66 FR 37958; (4) most of the scientific assessment work has been completed while meeting other Agency requirements, and only a modest additional effort will be needed to complete the review and documentation for IRIS.

The following IRIS health assessments have recently begun or will be started in FY 2002, with completion expected in FY 2003 or FY 2004, unless otherwise noted. It is for these substances that the Agency is primarily requesting information from the public for consideration in the assessments.

Unless otherwise noted, noncancer and cancer endpoints will be assessed for each substance. Pesticides denoted with an asterisk (*) will have only oral reference dose and carcinogenicity endpoints assessed. Substances denoted with a double asterisk (**) will be evaluated for effects from acute and subchronic exposure, in addition to chronic exposure. These substances constitute a pilot test to evaluate the application of methods, procedures, and resource needs for adding less-thanlifetime information to IRIS. For all endpoints assessed, both qualitative and quantitative assessments are being developed where information is available.

Name	CAS No.
Atrazine*	1912–24–9
Captan*	133-06-2
Di(2-ethylhexyl)adipate (DEHA)	103-23-1
Dibutyl phthalate	84–74–2
Ethalfluralin*	55283-68-6
gamma-Hexachlorocyclohexane	
(Lindane)*	58-89-9
Hydrogen cyanide †	74–90–8
Methyl ethyl ketone	78-93-3
2-Methylnaphthalene	91–57–6
Methomyl*	16752-77-5
Naphthalene (cancer; inh.	
route)	91–20–3
PAH mixtures	N.A.
Perfluorooctanoic acid—ammo-	
nium salt**	3825–26–1
Perfluorooctane sulfonate—po-	0020 20 1
tassium salt**	2795–39–3
Propachlor*	1918–16–7
Thallium†	7440–28–0
1,1,1-Trichloroethane**	71–55–6
	71 33 0

† = completion expected FY 2004/FY 2005.

Submission of Information

As in previous Federal Register notices announcing the annual IRIS agenda, EPA is soliciting public involvement in new assessments starting in FY 2002. While EPA conducts a thorough literature search for each chemical substance, there may be unpublished studies or other primary technical sources that we may not otherwise obtain through open literature searches. We would greatly appreciate receiving scientific information from the public during the information gathering stage for the list of "new assessments" listed above. Interested persons should provide scientific analyses, studies, and other pertinent scientific information. Also note that if you have submitted certain information previously, then there is no need to resubmit that information. While EPA is primarily soliciting information on new assessments announced in this notice, the public may submit information on any chemical substance at any time.

Procedures for Submission

Similar to the process described in the February 22, 2001, **Federal Register**, submissions will be handled in a threestep process:

1. Submission Inventory: First, you should simply provide a list within 60 days of this notice briefly identifying all the information (studies, reports, articles, etc.) you wish to submit. The list should specify by name and CASRN (Chemical Abstract Service Registry Number) the chemical substance(s) to which the information pertains, state the type of assessment that is being addressed (e.g., carcinogenicity), and describe briefly the information to be submitted for consideration. Where possible, documents should be listed in scientific citation format, that is, author(s), title, journal, and date. Your cover letter should state that the correspondence is an IRIS submission, describe in general terms the purpose of the submission, and include names, addresses, and telephone numbers of person(s) to contact for additional information. Mail two copies of the submission inventory to the IRIS Submission Desk, c/o ASRC, 6301 Ivy Lane, Suite 300, Greenbelt, MD 20770. Note that this is a new address for the Submission Desk.

Alternatively, you may submit the submission inventory and cover letter electronically to *IRIS.desk@epa.gov*. Electronic information must be submitted in WordPerfect format or as an ASCII file. Information also will be accepted on 3.5" floppy disks. All information in electronic form must be identified as an IRIS submission.

2. EPA Replies to Submission Inventory: In the second step, EPA will compare the submission inventory to existing files and identify the information that should be submitted. This step will help prevent an influx of duplicative information. You will receive notification of whether full submission of the information is requested.

3. Full Submission of Selected *Material:* In the third step, you should send in the information indicated by EPA within 30 days of EPA's reply. Prompt response to EPA will ensure that your material can be considered in the assessment in a timely fashion. Submissions should include a cover letter addressing all of the points in Item 1 above. In addition, when you submit results of new health effects studies concerning existing substances on IRIS, you should include a specific explanation of how and why the study results could change the information in IRIS.

Please send two copies, at least one of which should be unbound, to the IRIS Submission Desk, as described in Item 1. The IRIS Submission Desk will acknowledge receipt of your information.

Confidential Business Information (CBI) should not be submitted to the IRIS Submission Desk. CBI must be submitted to the appropriate EPA Office via established procedures for submission of CBI (see 40 CFR, Part 2, Subpart B). If you believe that a CBI submission contains information with implications for IRIS, please note that in the cover letter accompanying the submission to the appropriate office.

You may also request to augment your submission with a scientific briefing to EPA staff. Such requests should be made directly to Amy Mills, IRIS Program Director (see FOR FURTHER INFORMATION).

Dated: January 2, 2002.

Art Payne,

Acting Deputy Director, National Center for Environmental Assessment.

[FR Doc. 02–511 Filed 1–8–02; 8:45 am] **BILLING CODE 6560–50–P**

ENVIRONMENTAL PROTECTION AGENCY

[FRL-7127-1]

Clean Air Act Advisory Committee: Accident Prevention Subcommittee Meeting

AGENCY: Environmental Protection Agency (EPA).

ACTION: Notice of meeting.

SUMMARY: The Clean Air Act section 112(r) required EPA to publish regulations to prevent accidental releases of chemicals and to reduce the severity of those releases that do occur. These accidental release prevention requirements build on the chemical safety work begun by the Emergency Planning and Community Right-to-Know Act (EPCRA) which sets forth requirements for industry, State and local governments. On June 20, 1996, EPA published the final rule for risk management programs to address prevention of accidental releases. Facilities that are subject to the rule are required to implement a risk management program at their facility, and submit a summary of this information (the Risk Management Plan, RMP) to EPA. Approximately 15,000 RMPs have been submitted to EPA.

The Accident Prevention Subcommittee was created in September 1996 to advise EPA's Chemical Emergency Preparedness and Prevention Office (CEPPO) on these chemical accident prevention issues, specifically, section 112(r) of the Clean Air Act.

DATES: The Accident Prevention Subcommittee of the Clean Air Act Advisory Committee will hold a public meeting on January 24, 2002 from 8:30 a.m. to 3:00 p.m.

ADDRESSES: The meeting will be held at the Environmental Protection Agency, Ariel Rios North, Emergency Operations Center, Room B444, 1200 Pennsylvania Avenue, NW, Washington DC. Members of the public are welcome to attend in person.

FOR FURTHER INFORMATION CONTACT:

Members of the public desiring additional information about this meeting, should contact William Finan, Designated Federal Official, U.S. EPA (5104A), 1200 Pennsylvania Avenue, NW, Washington DC 20460, via the Internet at: finan.bill@epa.gov, by telephone at (202) 564–7981 or FAX at (202) 564–8444.

SUPPLEMENTARY INFORMATION:

Agenda

Opening Remarks—Jim Makris (8:30–9:00)

Update on Risk Management Plans Submitted to Date (9:00–9:45) Proposed Third-Party Audits for the RMP Program (9:45–10:45) National Chemical Safety Assessment (10:45–11:30)

Options on improving site security at chemical facilities (e.g., inclusion in company safety, health, and environment programs; legislation, regulation, standards, voluntary programs) (12:30–1:30)

How we can use upcoming Organization for Economic Cooperation and Development (OECD) publications in the US (the OECD publications are: Guidance on Safety Performance Indicators; and Guiding Principles for Chemical Accident Prevention, Preparedness, and Response (1:30—2:30) Comments from the public (2:30–3:00)

Members of the public who wish to make a brief oral presentation in person in Washington DC to the Subcommittee at the January 24 meeting must contact William Finan in writing (by letter, fax, or email—see previously stated information) no later than January 21, 2002 in order to be included on the agenda. Written comments may be submitted to the Accident Prevention Subcommittee up through the date of the meeting. Please address such material to William Finan at the above address.

The Accident Prevention
Subcommittee expects that public
statements presented at its meetings will
not be repetitive or previously
submitted oral or written statements. In
general, opportunities for oral comment
will be limited to no more than three
minutes per speaker and no more than
thirty minutes total. Written comments
(twelve copies) received sufficiently
prior to a meeting date (usually one
week prior to a meeting or
teleconference), may be mailed to the
Subcommittee prior to its meeting.

Additional information on the Accident Prevention Subcommittee is available on the Internet at: http://www.epa.gov/swercepp/acc-pre.html.

If you would like to automatically receive future information on the Accident Prevention Subcommittee and its Workgroups by email, you can subscribe to the EPA-CEPPO Listserve by following directions at www.epa.gov/ceppo.

Dated: January 3, 2002.

William Finan,

Designated Federal Official.
[FR Doc. 02–510 Filed 1–8–02; 8:45 am]

BILLING CODE 6560-50-P

ENVIRONMENTAL PROTECTION AGENCY

[OPP-64062; FRL-6815-9]

Notice of Receipt of Requests for Amendments to Delete Uses in Certain Pesticide Registrations

AGENCY: Environmental Protection Agency (EPA).

ACTION: Notice.

SUMMARY: In accordance with section 6(f)(1) of the Federal Insecticide, Fungicide and Rodenticide Act (FIFRA), as amended, EPA is issuing a notice of receipt of request for amendment by registrants to delete uses in certain pesticide registrations.

DATES: Unless a request is withdrawn, the Agency will approve these use deletions and the deletions will become effective on July 8, 2002, unless indicated otherwise.

FOR FURTHER INFORMATION CONTACT: By mail: James A. Hollins, Office of Pesticide Programs (7502C), Environmental Protection Agency, Ariel Rios Building, 1200 Pennsylvania Avenue, N.W., Washington, DC 20460. Office location for commercial courier delivery, telephone number and e-mail address: Rm. 266A, Crystal Mall #2, 1921 Jefferson Davis Highway, Arlington, VA 22202, (703) 305–5761; e-mail: hollins.james@epa.gov.