III. Existing Stocks Provisions

The Agency has authorized registrants to sell or distribute product under the previously approved labeling for a period of 18 months after approval of the revision, unless other restrictions have been imposed, as in special review actions.

List of Subjects

Environmental protection, Pesticides and pest, Product registrations.

Dated: November 3, 1995.

Frank Sanders,

Director, Program Management and Support Division, Office of Pesticide Programs.

[FR Doc. 95-28393 Filed 11-16-95; 8:45 am]

BILLING CODE 6560-50-F

[OPP-66219; FRL-4987-1]

Notice of Receipt of Requests to Voluntarily Cancel 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 requests by registrants to voluntarily cancel certain pesticide registrations.

DATES: Unless a request is withdrawn by Febuary 15, 1996, orders will be issued canceling all of these registrations.

FOR FURTHER INFORMATION CONTACT: By mail: James A. Hollins, Office of Pesticide Programs (7502C), Environmental Protection Agency, 401 M St., SW., Washington, DC 20460. Office location for commercial courier delivery and telephone number: Room 216, Crystal Mall No. 2, 1921 Jefferson

Davis Highway, Arlington, VA, (703) 305–5761; e-mail: hollins.james@epamail.epa.gov.

SUPPLEMENTARY INFORMATION:

I. Introduction

Section 6(f)(1) of the Federal Insecticide, Fungicide and Rodenticide Act (FIFRA), as amended, provides that a pesticide registrant may, at any time, request that any of its pesticide registrations be canceled. The Act further provides that EPA must publish a notice of receipt of any such request in the Federal Register before acting on the request.

II. Intent to Cancel

This Notice announces receipt by the Agency of requests to cancel some 25 pesticide products registered under section 3 or 24(c) of FIFRA. These registrations are listed in sequence by registration number (or company number and 24(c) number) in the following Table 1.

TABLE 1--Registrations With Pending Requests for Cancellation

Registration no.	Product Name	Chemical Name
000299-00221	Martin's Ear - Tix - Tox Ear Tick Control	O,O-Diethyl O-(3,5,6-trichloro-2-pyridyl) phosphorothioate
000352-00396	Dupont Benlate DF Fungicide	Methyl 1-(butylcarbamoyl)-2-benzimidazolecarbamate
000655-00790	Prentox Larva-Lur	Dimethyl (2,2,2-trichloro-1-hydroxyethyl)phosphonate
002393 OR-86-0008	Mylone 99G Soil Fumigant NC	Tetrahydro-3,5-dimethyl-2 <i>H</i> -1,3,5-thiadiazine-2-thione
002935-00416	Wilbur Ellis MCPA Sodium Salt.	Sodium 2-methyl-4-chlorophenoxyacetate
003125 CA-82-0103	Meta-Systox-R Spray Concentrate	S-(2-(Ethylsulfinyl)ethyl) O,O-dimethyl phosphorothioate
003125 CA-82-0104	Meta-Systox-R Spray Concentrate	S-(2-(Ethylsulfinyl)ethyl) O,O-dimethyl phosphorothioate
003125 CA-91-0029	Metasystox-R Spray Concentrate	S-(2-(Ethylsulfinyl)ethyl) O,O-dimethyl phosphorothioate
003125 ID-77-0006	Meta-Systox-R Spray Concentrate	S-(2-(Ethylsulfinyl)ethyl) O,O-dimethyl phosphorothioate
003125 ID-78-0001	Meta-Systox-R Spray Concentrate	S-(2-(Ethylsulfinyl)ethyl) O,O-dimethyl phosphorothioate
003125 NM-79-0025 .	Furadan 4 Flowable	2,3-Dihydro-2,2-dimethyl-7-benzofuranyl methylcarbamate
N/A	N/A	2,3-Dihydro-2,2-dimethyl-7-benzofuranyl methylcarbamate
N/A	N/A	S-(2-(Ethylsulfinyl)ethyl) O,O-dimethyl phosphorothioate
003125 NV-77-0014	Meta-Systox-R Spray Concentrate	S-(2-(Ethylsulfinyl)ethyl) O,O-dimethyl phosphorothioate
003125 OH-89-0004	Metasystox-R Spray Concentrate	S-(2-(Ethylsulfinyl)ethyl) O,O-dimethyl phosphorothioate
003125 OR-77-0019	Meta-Systox-R Spray Concentrate	S-(2-(Ethylsulfinyl)ethyl) O,O-dimethyl phosphorothioate
003125 OR-79-0067	Meta-Systox-R Spray Concentrate	S-(2-(Ethylsulfinyl)ethyl) O,O-dimethyl phosphorothioate
003125 WA-77-0014 .	Meta-Systox-R Spray Concentrate	S-(2-(Ethylsulfinyl)ethyl) O,O-dimethyl phosphorothioate
003125 WA-77-0058 .	Meta-Systox-R Spray Concentrate	S-(2-(Ethylsulfinyl)ethyl) O,O-dimethyl phosphorothioate
003125 WA-80-0077 .	Meta-Systox-R Spray Concentrate	S-(2-(Ethylsulfinyl)ethyl) O,O-dimethyl phosphorothioate
003125 WA-85-0003 .	Meta-Systox-R Spray Concentrate	S-(2-(Ethylsulfinyl)ethyl) O,O-dimethyl phosphorothioate
003125 WA-89-0033 .	Metasystox-R Spray Concentrate	S-(2-(Ethylsulfinyl)ethyl) O,O-dimethyl phosphorothioate
007056-00149	CSA Screwworm Spray	Dipropyl isocinchomeronate
N/A	N/A	O,O-Diethyl O-(3,5,6-trichloro-2-pyridyl) phosphorothioate
010182 OR-88-0016	Fusilade 2000 Herbicide	Butyl (R)-2-(4-((5-(trifluoromethyl)-2-pyridinyl)oxy)phenoxy)propanoate
010182 OR-90-0018	Fusilade 2000 Herbicide	Butyl (R)-2-(4-((5-(trifluoromethyl)-2-pyridinyl)oxy)phenoxy)propanoate
050534-00028	75% Dimethyl T	Dimethyl tetrachloroterephthalate

Unless a request is withdrawn by the registrant within 90 days of publication of this notice, orders will be issued canceling all of these registrations. Users of these pesticides or anyone else desiring the retention of a registration should contact the applicable registrant directly during this 90-day period.

The following Table 2 includes the names and addresses of record for all

registrants of the products in Table 1, in sequence by EPA Company Number.

TABLE 2--Registrants Requesting Voluntary Cancellation

EPA Com- pany no.	Company Name and Address
000299 .	C. J. Martin Co, Box 630009, Nacogdoches, TX 75963.

TABLE 2--Registrants Requesting Voluntary Cancellation—Continued

EPA Com- pany no.	Company Name and Address
000352 .	E. I. Du Pont De Nemours & Co, Inc., Barley Mill Plaza, Walker's Mill, Wilmington, DE 19880.
000655 .	Prentiss Inc., 21 Vernon Street, C.B. 2000, Floral Park, NY 11001.
002393 .	Haco, Inc., Box 7190, Madison, WI 53707.
002935 .	Wilbur Ellis Co., 191 W. Shaw Ave, Fresno, CA 93704.
003125 .	Bayer Corp., Agriculture Division, 8400 Hawthorn Rd., Box 4913, Kansas City, MO 64120.
007056 .	IQ Products Co, Attn: Marty York, 16212 State Hwy 249, Houston, TX 77086.
010182 .	Zeneca Ag Products, Box 15458, Wilmington, DE 19850.
050534 .	ISK Biosciences Corp., 5966 Heisley Rd., Box 8000, Mentor, OH 44061.

III. Procedures for withdrawal of request

Registrants who choose to withdraw a request for cancellation must submit such withdrawal in writing to James A. Hollins, at the address given above, postmarked before February 15, 1996. This written withdrawal of the request for cancellation will apply only to the applicable 6(f)(1) request listed in this notice. If the product(s) have been subject to a previous cancellation action, the effective date of cancellation and all other provisions of any earlier cancellation action are controlling. The withdrawal request must also include a commitment to pay any reregistration fees due, and to fulfill any applicable unsatisfied data requirements.

IV. Provisions for Disposition of Existing Stocks

The effective date of cancellation will be the date of the cancellation order. The orders effecting these requested cancellations will generally permit a registrant to sell or distribute existing stocks for one year after the date the cancellation request was received. This policy is in accordance with the Agency's statement of policy as prescribed in Federal Register No. 123, Vol. 56, dated June 26, 1991. Exceptions to this general rule will be made if a product poses a risk concern, or is in noncompliance with reregistration requirements, or is subject to a data callin. In all cases, product-specific disposition dates will be given in the cancellation orders.

Existing stocks are those stocks of registered pesticide productswhich are currently in the United States and which have beenpackaged, labeled, and released for shipment prior to the effectivedate of the cancellation action. Unless the provisions of anearlier order apply, existing stocks already in the hands ofdealers or users can be distributed, sold or used legally untilthey are exhausted, provided that such further sale and use comply with the EPA-approved label and labeling of the affected product(s). Exceptions to these general rules will be made in specific cases when more stringent restrictions on sale, distribution, or use of the products or their ingredients have already been imposed, as in Special Review actions, or where the Agency has identified significant potential risk concerns associated with a particular chemical.

List of Subjects

Environmental protection, Pesticides and pest, Product registrations.

Dated: November 3, 1995.

Frank Sanders,

Director, Program Management & Support Division, Office of Pesticide Programs.

[FR Doc. 95–28392 Filed 11–16–95; 8:45 am] BILLING CODE 6560–50–F

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. 94N-0376]

Plascon, Inc.; Opportunity for Hearing on a Proposal to Revoke U.S. License No. 572–003

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing an opportunity for a hearing on a proposal to revoke the establishment license (U.S. License No. 572–003) and the product license issued to Plascon, Inc., doing business as Anderson Plasma Center, for the manufacture of Source Plasma. The proposed revocation is based on the firm's history of continued noncompliance with the applicable biologics regulations and the license standards.

DATES: The firm may submit a written request for a hearing to the Dockets Management Branch by December 18, 1995, and any data and information justifying a hearing by January 16, 1996.

Other interested persons may submit written comments on the proposed revocation by January 16, 1996.

ADDRESSES: Submit written requests for a hearing, any data and information justifying a hearing, and any comments on the proposed revocation to the Dockets Management Branch (HFA–305), Food and Drug Administration, rm. 1–23, 12420 Parklawn Dr., Rockville, MD 20857.

FOR FURTHER INFORMATION CONTACT: Timothy W. Beth, Center for Biologics Evaluation and Research (HFM–635), Food and Drug Administration, 1401 Rockville Pike, Rockville, MD 20852–1448, 301–594–3074.

SUPPLEMENTARY INFORMATION: FDA is proposing to revoke the establishment license (U.S. License No. 572–003) and the product license issued to Plascon, Inc., doing business as Anderson Plasma Center, 2507 Nichol Ave., Anderson, IN 46011, for the manufacture of Source Plasma.

During preapproval and routine inspections conducted by FDA at Plascon, Inc., in 1989, 1991, 1992, and 1993, significant deviations from the applicable Federal regulations and the license standards were documented. Following each of these inspections, FDA provided to Plascon, Inc., written documentation of the deviations observed; FDA then requested that Plascon, Inc., indicate in writing what corrective action plan would be undertaken to remedy the deviations. Following the 1992 inspection, FDA issued a warning letter dated November 12, 1992, to Plascon, Inc., advising the firm that failure to promptly correct the deviations observed during the inspection could result in regulatory action by FDA without further notice.

In response to FDA's inspectional observations, in letters dated December 19, 1989, October 17, 1991, and January 6, 1993, Plascon, Inc., proposed corrective action plans. However, subsequent inspections of Plascon, Inc., by FDA continued to demonstrate that sufficient and effective long-term corrective action had not been achieved. Plascon, Inc.'s, cumulative inspectional history thus established a pattern of continued noncompliance with the applicable Federal regulations and license standards.

The most recent inspection, conducted from December 13 through December 17, 1993, revealed continuing significant deviations from the applicable regulations and the license standards. These deviations included, but were not limited to, the following: (1) Failure to adequately determine donor suitability (21 CFR 606.100(b)(1)