notice of availability of the agency's finding of no significant impact and the evidence supporting that finding will be published with the regulation in the **Federal Register** in accordance with 21 CFR 25.40(c).

Dated: June 23, 1995.

Alan M. Rulis,

Acting Director, Office of Premarket Approval, Center for Food Safety and Applied Nutrition.

[FR Doc. 95–17093 Filed 7–11–95; 8:45 am] BILLING CODE 4160–01–F

[Docket No. 93F-0244]

National Starch and Chemical Co.; Withdrawal of Food Additive Petition

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing the withdrawal, without prejudice to a future filing, of a food additive petition (FAP 3B4385) proposing that the food additive regulations be amended to provide for the safe use of starch, modified by treatment with diethylaminoethylchloride, hydrochloride salt and 2-chloro-*N*-(2,2-dimethoxyethyl)-*N*-methylacetamide as an internal sizing for paper and paperboard intended to contact food.

FOR FURTHER INFORMATION CONTACT: Mitchell A. Cheeseman, Center for Food Safety and Applied Nutrition (HFS–217), Food and Drug Administration, 200 C St. SW., Washington, DC 20204, 202–418–3083.

SUPPLEMENTARY INFORMATION: In a notice published in the Federal Register of August 12, 1993 (58 FR 42977), FDA announced that a food additive petition (FAP 3B4385) had been filed by National Starch and Chemical Co., Finderne Ave., Bridgewater, NJ 08807. The petition proposed to amend the food additive regulations in § 178.3520 Industrial starch-modified (21 CFR 178.3520) to provide for the safe use of starch, modified by treatment with diethylaminoethylchloride, hydrochloride salt (CAS Reg. No. 869– 24-9) and 2-chloro-N-(2,2dimethoxyethyl)-N-methylacetamide (CAS Reg. No. 69184-36-7) as an internal sizing for paper and paperboard intended to contact food. National Starch and Chemical Co. has now withdrawn the petition without prejudice to a future filing (21 CFR 171.7).

Dated: June 23, 1995.

Alan M. Rulis,

Acting Director, Office of Premarket Approval, Center for Food Safety and Applied Nutrition.

[FR Doc. 95–17097 Filed 7–11–95; 8:45 am] BILLING CODE 4160–01–F

[Docket No. 89F-0111]

Rhone-Poulenc, Inc.; Withdrawal of Food Additive Petition

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing the withdrawal, without prejudice to future filing, of a food additive petition (FAP 9B4140) proposing that the food additive regulations be amended to provide for the safe use of alkyl (C_{14} – C_{30}) benzene as a component of adhesives for articles intended to contact food.

FOR FURTHER INFORMATION CONTACT: Hortense S. Macon, Center for Food Safety and Applied Nutrition (HFS–216), Food and Drug Administration, 200 C St. SW., Washington, DC 20204, 202–418–3086.

SUPPLEMENTARY INFORMATION: In a notice published in the Federal Register of May 2, 1989 (54 FR 18700), FDA announced that a food additive petition (FAP 9B4140) had been filed by Rhone-Poulenc, Inc., 1669 Corporate Rd. West, Lakewood, NJ 08071. The petition proposed to amend the food additive regulations in § 175.105 Adhesives (21 CFR 175.105) to provide for the safe use of alkyl (C₁₄-C₃₀) benzene as a component of adhesives for articles intended to contact food. Rhone-Poulenc, Inc., has now withdrawn the petition without prejudice to a future filing (21 CFR 171.7).

Dated: June 23, 1995.

Alan M. Rulis,

Acting Director, Office of Premarket Approval, Center for Food Safety and Applied Nutrition.

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Health Care Financing Administration

Statement of Organization, Functions, and Delegations of Authority; Office of the Actuary

Part F of the Statement of Organization, Functions, and Delegations of Authority for the Department of Health and Human Services, Health Care Financing Administration (HCFA), (**Federal Register**, Vol. 59, No. 60, pp. 14643–14644, dated Tuesday, March 29, 1994) is amended to reflect changes in the substructure of the Office of the Actuary (OACT). The OACT functional statement has not been changed; however, the remaining OACT substructure is being published to reflect the organizational changes resulting from streamlining efforts.

The specific amendments to Part F are as follows:

- Section F.10.C.4. (Organization) is amended to read as follows:
 - 4. Office of the Actuary
- a. Office of Medicare and Medicaid Cost Estimates
 - b. Office of National Health Statistics
- Section F.20.C.4. (Functions) is amended by modifying office statements and deleting the office substructure in their entirety. The new functional statements read as follows:

a. Office of Medicare and Medicaid Cost Estimates (FKC1)

- Prepares cost estimates for the Hospital Insurance (HI) program, the Supplementary Medical Insurance (SMI) program, and the Medicaid program for use in the President's budget.
- Evaluates the operations of the Medicare trust funds particularly relating to outlays and program solvency.
- Develops such variables as the Part B premium rates, the inpatient hospital deductible, the Part A premium rate for voluntary enrollees, and the physicians' economic index applicable to prevailing fees
- Develops the payment rates for the annual update of the adjusted average per capita cost (AAPCC) ratebook, which is used to pay health maintenance organizations that enter into a risk contract with HCFA to provide benefits to Medicare enrollees.
- Provides cost estimates for the Medicaid program, including the development of cost estimates for proposed changes in Medicaid or in programs affecting Medicaid, and overall Medicaid program costs for years after the current budget year.
- Serves as technical consultant throughout the Government on Medicare and Medicaid cost estimate issues.
- Provides actuarial consultation to other organizations in the research of AAPCC methodology.

b. Office of National Health Statistics (FKC2)

• Develops, maintains and makes analytical use of the National Health Accounts (NHA) which include annual