

The Director, Management Analysis and Services Office, has been delegated the authority to sign **Federal Register** notices pertaining to announcements of meetings and other committee management activities for both the CDC and ATSDR.

Dated: January 10, 2006.

**Alvin Hall,**

*Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.*

[FR Doc. E6-529 Filed 1-18-06; 8:45 am]

**BILLING CODE 4163-18-P**

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Food and Drug Administration

[Docket No. 2006N-0010]

#### **Able Laboratories, Inc.; Withdrawal of Approval of 43 Abbreviated New Drug Applications**

**AGENCY:** Food and Drug Administration, HHS.

**ACTION:** Notice.

**SUMMARY:** The Food and Drug Administration (FDA) is withdrawing approval of 43 abbreviated new drug applications (ANDAs) held by Able Laboratories, Inc. (Able Labs), One Able Dr., Cranbury, NJ 08512. The drug

products are no longer marketed, and Able Labs has requested that the approval of the applications be withdrawn.

**DATES:** Effective January 19, 2006.

**FOR FURTHER INFORMATION CONTACT:** Florine P. Purdie, Center for Drug Evaluation and Research (HFD-7), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301-594-2041.

**SUPPLEMENTARY INFORMATION:** The applications listed in the table in this document are no longer marketed, and Able Labs has requested that FDA withdraw approval of the applications. The company has also, by its request, waived its opportunity for a hearing.

Application No.	Drug
40-390	Butalbital, Acetaminophen, and Caffeine Tablets USP, 50 milligrams (mg)/325 mg/40 mg
40-394	Butalbital, Acetaminophen, and Caffeine Tablets USP, 50 mg/500 mg/40 mg
40-402	Phentermine Hydrochloride (HCl) Tablets USP, 37.5 mg
40-403	Phentermine HCL Capsules USP, 30 mg (powder)
40-413	Methocarbamol Tablets USP, 500 mg and 750 mg
40-421	Carisoprodol Tablets USP, 350 mg
40-427	Phentermine HCl Capsules USP, 30 mg (beads)
40-449	Promethazine HCl Suppositories USP, 50 mg
40-464	Hydrocodone Bitartrate and Acetaminophen Tablets USP, 7.5 mg/325 mg and 10 mg/325 mg
40-469	Hydrocodone Bitartrate and Acetaminophen Tablets USP, 7.5 mg/750 mg
40-473	Hydrocodone Bitartrate and Acetaminophen Tablets USP, 10 mg/500 mg
40-474	Hydrocodone Bitartrate and Acetaminophen Tablets USP, 7.5 mg/650 mg
40-476	Hydrocodone Bitartrate and Acetaminophen Tablets USP, 10 mg/650 mg
40-477	Hydrocodone Bitartrate and Acetaminophen Tablets USP, 5 mg/500 mg
40-478	Hydrocodone Bitartrate and Acetaminophen Tablets USP, 5 mg/325 mg
40-483	Bethanechol Chloride Tablets USP, 10 mg
40-485	Bethanechol Chloride Tablets USP, 25 mg
40-490	Hydrocodone Bitartrate and Acetaminophen Tablets USP, 7.5 mg/500 mg
40-492	Bethanechol Chloride Tablets USP, 5 mg
40-497	Phentermine HCl Capsules USP, 15 mg
40-504	Promethazine HCl Suppositories USP, 12.5 mg and 25 mg
40-509	Bethanechol Chloride Tablets USP, 50 mg
40-529	Methamphetamine HCl Tablets USP, 5 mg
40-539	Theophylline Extended-Release Tablets, 600 mg
40-543	Theophylline Extended-Release Tablets, 400 mg
40-546	Theophylline Extended-Release Tablets, 450 mg

Application No.	Drug
40-548	Theophylline Extended-Release Tablets, 300 mg
40-558	Promethazine HCl Tablets USP, 12.5 mg, 25 mg, and 50 mg
40-559	Hydroxyzine HCl Tablets USP, 10 mg
40-562	Hydroxyzine HCl Tablets USP, 25 mg
40-563	Hydroxyzine HCl Tablets USP, 50 mg
76-114	Indomethacin Extended-Release Capsules USP, 75 mg
76-121	Lithium Carbonate Capsules USP, 300 mg
76-382	Lithium Carabonate Extended-Release Tablets USP, 300 mg
76-462	Metronidazole Extended-Release Tablets, 750 mg
76-505	Metronidazole Capsules, 375 mg
76-519	Metronidazole Tablets USP, 250 mg and 500 mg
76-528	Butalbital, Acetaminophen, Caffeine, and Codeine Phosphate Capsules, 50 mg/325 mg/40 mg/30 mg
76-544	Naproxen Sodium Tablets USP, 275 mg and 550 mg
76-666	Indomethacin Capsules USP, 25 mg and 50 mg
76-814	Dextroamphetamine Sulfate Extended-Release Capsules, 5 mg, 10 mg, and 15 mg
76-823	Lithium Carbonate Capsules USP, 150 mg, 300 mg, and 600 mg
76-907	Atenolol Tablets USP, 25 mg, 50 mg, and 100 mg

Therefore, under section 505(e) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 355(e)) and under authority delegated to the Director, Center for Drug Evaluation and Research, by the Commissioner of Food and Drugs, approval of the applications listed in the table in this document, and all amendments and supplements thereto, is hereby withdrawn, effective January 19, 2006.

Dated: January 4, 2006.

**Douglas C. Throckmorton,**

*Deputy Director, Center for Drug Evaluation and Research.*

[FR Doc. E6-506 Filed 1-18-06; 8:45 am]

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## DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Indian Health Service

#### Request for Public Comment: 60-Day Proposed Information Collection: Indian Health Service Chief Executive Officer Retention Survey

**AGENCY:** Indian Health Service, HHS.

**SUMMARY:** The Department of Health and Human Services, as part of its continuing effort to reduce paperwork and respondent burden, conducts a pre-clearance consultation program to

provide the general public and Federal agencies with an opportunity to comment on proposed and/or continuing collections of information in accordance with the Paperwork Reduction Act of 1995 (PRA95) (44 U.S.C. 3506(c)(2)(A)). This program helps to ensure that requested data can be provided in the desired format, reporting burden (time and financial resources) is minimized, collection instruments are clearly understood, and the impact of collection requirements on respondents can be properly assessed. Currently, the Indian Health Service (IHS) is providing a 60-day advance opportunity for public comment on a proposed extension of current information collection activity to be submitted to the Office of Management and Budget for review.

#### Proposed Collection

*Title:* 0917-NEW, "Indian Health Service Chief Executive Officer Retention Survey".

*Type of Information Collection*

*Request:* New Collection.

*Form Number:* None.

*Forms:* The IHS Chief Executive Officer Retention Survey.

*Need and Use of Information*

*Collection:* The National Council of Chief Executive Officers (NCCEO) was established to ensure that the IHS

Service Unit Chief Executive Officers (CEO) effectively participate in the establishment and implementation of strategies to achieve the IHS mission. Part of their responsibility (as stated in their Charter) includes: ongoing recruitment, development, and retention of professional CEOs. The NCCEO's purpose is to ensure that the IHS Service Unit CEO and their Tribal CEO counterparts effectively participate in the establishment and implementation of an agency strategy to achieve the IHS mission. The current Executive Committee is actively addressing recruitment, retention and succession planning for their constituents, the IHS CEOs. To enhance their ability to be effective in this challenging task, the NCCEO needs to know more about IHS CEOs and the issues that affect retention and recruitment including the competitive influences of private sector health care delivery systems. The chosen method to obtain this critical information from the CEOs of IHS, Tribal and Urban facilities is by electronic survey. The goal of the IHS is to raise the health status of American Indians and Alaska Natives to the highest possible level. To meet this goal, the IHS is committed to providing high quality health services to the eligible service population. An important factor