## DEPARTMENT OF HEALTH & HUMAN SERVICES

Public Health Service

Food and Drug Administration Rockville, MD 20857

NDA 11-522/S-040

Duramed Research Inc. Attention: Joseph A. Carrado, M.Sc., R.Ph. Vice President, Clinical Regulatory Affairs One Belmont Ave 11th Floor Bala Cynwyd, PA 190047

Dear Mr. Carrado:

We acknowledge receipt of your supplemental new drug application dated April 16, 2007, received April 17, 2007, submitted under section 505(b) of the Federal Food, Drug, and Cosmetic Act for Adderall (mixed salts of a single-entity amphetamine product) Tablets.

Reference is also made to an Agency letter dated February 21, 2006 and March 19, 2007 (electronic communication), requesting Medication Guides for all CNS stimulant products to treat Attention-Deficit Hyperactivity Disorder (ADHD).

Your April 16, 2007 submission provides for a response to our February 21, 2006 and March 19, 2007 (electronic communication) action letter.

This supplement, submitted under "Changes Being Effected", provides for revisions to the "PRECAUTIONS-Information for Patients," section, a revised package insert incorporating the new Medication Guide in place of the patient information leaflet, and revised container labeling for each strength of Adderall instructing the authorized dispenser to provide a Medication Guide to each patient that receives a prescription for the product.

We have completed our review of this application and it is approved, effective on the date of this letter, for use as recommended in the enclosed labeling text.

If you issue a letter communicating important information about this drug product (i.e., a "Dear Health Care Professional" letter), we request that you submit a copy of the letter to this NDA and a copy to the following address:

MEDWATCH Food and Drug Administration 5515 Security Lane HFD-001, Suite 5100 Rockville, MD 20852 NDA 11-522/S-040 Page 2

We remind you that you must comply with reporting requirements for an approved NDA (21 CFR 314.80 and 314.81).

If you have any questions, contact LT Felecia Curtis, Regulatory Project Manager, at (301) 796-0877.

Sincerely,

{See appended electronic signature page}

Thomas Laughren, MD Director Division of Psychiatry Products Office of Drug Evaluation I Center for Drug Evaluation and Research

Enclosure

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this page is the manifestation of the electronic signature.	

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Thomas Laughren 6/7/2007 04:05:26 PM