Dear Dr. Lamendola:


We acknowledge receipt of your submissions dated February 12, March 22 and 28 (2), July 2, August 9 (2), September 6, 16, and 25, October 24, and November 12, 13, and 15, 2002.

These supplemental new drug applications provide for the over-the-counter use of Claritin (loratadine) Tablets, Syrup, and RediTabs for the temporary relief of symptoms of hay fever or other upper respiratory allergies: runny nose, sneezing, itchy, watery eyes, and itching of the nose or throat.

We have completed our review of these supplemental applications as amended. These supplemental applications are approved, effective on the date of this letter, for use as recommended in the agreed-upon labeling text.

The final printed labeling (FPL) must be identical to the submitted draft labeling (immediate container and carton labels, blister backing, launch pouch sticker, and pouch submitted November 12, 2002), and must be in the “Drug Facts” format (21 CFR 201.66). Marketing the products with FPL that is not identical to the approved labeling text and in the required format may render the products misbranded and unapproved new drugs.

Please submit an electronic version of the FPL according to the guidance for industry titled Providing Regulatory Submissions in Electronic Format - NDA. Alternatively, you may submit 20 paper copies of the FPL as soon as it is available but no more than 30 days after it is printed. Individually mount ten of the copies on heavy-weight paper or similar material. For administrative purposes, designate these submissions “FPL for approved NDAs 19658/S-018, 20-704/S-008, and 20-641/S-009.” Approval of this submission by FDA is not required before the labeling is used.
We remind you of your post-approval follow-up agreements in your submission dated November 13, 2002. These include submitting information in each quarterly periodic safety report for the first three years after approval on reports from various sources of the occurrence of cases of hypospadias. Also, in your periodic reports, you will separately analyze and present adverse events in the 2-5 year-old age group and the less-than-2 year-old age group.

Oversight of this application is being transferred to the Division of Over-the-Counter Drug Products (HFD-560). If you have any questions, call Elaine Abraham, R.Ph., Regulatory Management Officer, at (301) 827-2301.

Sincerely,

[See appended electronic signature page]  [See appended electronic signature page]

Jonca Bull, M.D.  Robert Meyer, M.D.
Director  Director
Office of Drug Evaluation V  Office of Drug Evaluation II
Center for Drug Evaluation and Research  Center for Drug Evaluation and Research
This is a representation of an electronic record that was signed electronically and this page is the manifestation of the electronic signature.

/s/

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Jonca Bull
11/27/02 08:22:45 AM

Robert Meyer
11/27/02 10:33:36 AM