



NDA 18-644/S-027/S-028
NDA 20-358/S-031/S-032
NDA 21-515/S-006/S-007

GlaxoSmithKline
Attention: Mary E. Martinson
Director, Psychiatry US Regulatory Affairs
Five Moore Drive, P.O. Box 13398
Research Triangle Park, NC 27709

Dear Ms. Martinson:

We acknowledge receipt of your supplemental new drug applications dated April 28, 2004 (NDAs 18-644/S-027, 20-358/S-031, & 21-515/S-006), and May 11, 2004 (NDAs 18-644/S-028, 20-358/S-032, & 21-515/S-007), submitted under section 505(b) of the Federal Food, Drug, and Cosmetic Act for Wellbutrin Immediate Release Tablets (NDA 18-644), Wellbutrin SR (bupropion hydrochloride) Sustained-Release Tablets (NDA 20-358), and Wellbutrin XL (bupropion hydrochloride) Extended-Release Tablets (NDA 21-515).

Supplemental applications 18-644/S-028, 20-358/S-032, & 21-515/S-007, submitted as "Changes Being Effected" submissions, provide for the following changes to product labeling as requested in our Agency letter dated March 19, 2004, and revised in an electronic communication to you from Paul David, of this office, on April 19, 2004:

1. The addition of a new subsection under **WARNINGS** entitled **Clinical Worsening and Suicide Risk**.
2. Revisions to the **PRECAUTIONS-Information for Patients** section.
3. Delete the section in **PRECAUTIONS-General** entitled "Suicide".
4. Add a reference to the **WARNINGS** section at the end of the **PRECAUTIONS- Pediatric Use** section, i.e., (see **WARNINGS-Clinical Worsening and Suicide Risk**).

We have completed the review of these supplemental applications and have concluded that adequate information has been presented to demonstrate that the drug product is safe and effective for use as recommended in your labeling submitted on May 11, 2004. Accordingly, these supplemental applications are approved effective on the date of this letter.

We note that your supplemental applications, 18-644/S-027, 20-358/S-031, & 21-515/S-006, were superseded by your May 11, 2004 submission. Therefore, we are going to administratively close these supplements and retain them in our files.

NDA 18-644/S-027/S-028, 20-358/S-031/S-032, & 21-515/S-006/S-007

Page 2

If a letter communicating important information about this drug product (i.e., a "Dear Health Care Practitioner" letter) is issued to physicians and others responsible for patient care, we request that you submit a copy of the letter to this NDA and a copy to the following address:

MEDWATCH, HF-2
FDA
5600 Fishers Lane
Rockville, MD 20857

We remind you that you must comply with the requirements for an approved NDA set forth under 21 CFR 314.80 and 314.81.

If you have any questions, call Mr. Paul David, R.Ph., Senior Regulatory Project Manager, at (301) 594-5530.

Sincerely,

{See appended electronic signature page}

Russell Katz, M.D.
Director
Division of Neuropharmacological Drug Products
Office of Drug Evaluation I
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/

Russell Katz
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