

Food and Drug Administration Rockville MD 20857

FILE COPY

December 2, 2013

Terri Nataline Principal Associate Lachman Consultant Services, Inc. 1600 Stewart Avenue, Suite 604 Westbury, NY 11590

Dear Ms. Nataline:

Your petition to the Food and Drug Administration requesting the Agency to determine whether Zovirax (acyclovir sodium) for injection, eq. 1 g base/vial (GlaxoSmithKline (GSK)), NDA 18 603 has been voluntarily withdrawn for safety or effectiveness reasons, was received by this office on 11/18/2013. It was assigned docket number FDA-2013-P-1515/CP1, and it was filed on 12/2/2013. Please refer to this docket number in future correspondence on this subject with the Agency.

Please note that the acceptance of the petition for filing is a procedural matter in that it in no way reflects an agency decision on the substantive merits of the petition.

Sincerely,

Karen Kennard

Director

Division of Dockets Management

Keren Kennard

FDA/Office of the Executive Secretariat (OES)