



**DEPARTMENT OF HEALTH & HUMAN SERVICES**

Public Health Service

Food and Drug Administration  
Silver Spring MD 20993

June 24, 2019

David Rosen  
Foley & Lardner LLP  
3000 K St., N.W., Suite 600  
Washington, DC 20007

Sent via email to: [drosen@foley.com](mailto:drosen@foley.com)

Dear Petitioner:

Your petition to the Commissioner of Food and Drug Administration requesting to declare that:

1. Sitagliptin Capsules Eq. 25 mg base, Eq. 50 mg base and Eq. 100 mg base are suitable for submission as an Abbreviated New Drug Application (ANDA), pursuant to 21 CFR §314.94; and
2. The reference product on which the contents of this petition are based is JANUVIA® (Sitagliptin) Tablets 25 mg, 50 mg and 100 mg;

Your submission was received by this office on 06/24/2019. It was assigned docket number FDA-2019-P-3024. 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,

Dynna Bigby  
Supervisory Administrative Proceedings Specialist  
Division of Dockets Management  
FDA/Office of Operations (OO)