



March 4, 2022

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*Sent via email to:* [Molly.Ventrelli@fresenius-kabi.com](mailto:Molly.Ventrelli@fresenius-kabi.com)

Dear Petitioner:

Your submission requesting that the Commissioner of Food and Drug designate Calcium Gluconate in Sodium Chloride Injection 1g/50mL and 2g/100mL manufactured by FK USA (NDA 208418), as a therapeutic equivalent, with an 'AP' rating, to the Calcium Gluconate in Sodium Chloride Injection 1g/50mL and 2g/100mL NDA 210906, by HQ Specialty Pharma Corp was received and processed under CFR 10.30 by this office on 03/04/2022.

It was assigned docket number FDA-2022-P-0290. Please refer to this docket number in future correspondence on this subject with the Agency.

Please note, the acceptance of the petition for filing is a procedural matter and in no way reflects the Agency's decision on the substantive merits of the petition.

Sincerely,

Dynna Bigby  
Supervisory Administrative Proceedings Officer  
Dockets Management Staff  
FDA/Office of Operations (OO)