



**Suitability Petition  
Completeness Assessment Correspondence**

Premier Research Consulting, LLC  
8000 Jarvis Avenue, Suite 100  
Newark, CA 94560  
Attn: Seth D. DePuy

Sent via email to: [seth.depuy@premierconsulting.com](mailto:seth.depuy@premierconsulting.com)

Docket No. FDA-2024-P-3573

Dear Seth D. DePuy:

This is in reference to your petition received on July 29, 2024, by the U.S. Food and Drug Administration (FDA or Agency), requesting permission, in accordance with 21 C.F.R. §314.93(b), to submit an Abbreviated New Drug Application (ANDA) for the following drug product: Lisinopril and Hydrochlorothiazide Oral Liquid, 10 mg/12.5 mg per 5 mL, 20 mg/12.5 mg per 5 mL, and 20 mg/25 mg per 5 mL. The petition was examined for initial completeness (i.e. Completeness Assessment) under CFR 10.30 and the Agency has made a threshold determination that the petition is complete.

This petition is subject to the provisions of the Generic Drugs User Fee Amendments of 2022 (GDUFA III) whereby any suitability petitions submitted in FY 2024-2027 will receive a goal date of 6 months after the completeness assessment. The GDUFA goal date for review of this petition is February 8, 2025.

If you have any questions, contact [ANDAFiling@fda.hhs.gov](mailto:ANDAFiling@fda.hhs.gov).

A copy of this letter will be placed on public display in the Dockets Management



Branch, Room 1061, Mail Stop HFA-305, 5630 Fishers Lane, Rockville, MD 20852.

Sincerely,

*{See appended electronic signature page}*

Diana Guon, Pharm.D., BCPS  
Pharmacist  
Division of Filing Review  
Office of Regulatory Operations  
Office of Generic Drugs  
Center for Drug Evaluation and Research



Diana  
Guon

Digitally signed by Diana Guon

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