

Food and Drug Administration Rockville MD 20857

MAR 2.0 2014

Edward J. Allera Buchanan Ingersoll & Rooney PC 1700 K Street N.W., Suite 300 Washington, DC 20006

Re:

Docket No. FDA-2013-P-1377

Dear Mr. Allera:

I am writing to inform you that the Food and Drug Administration (FDA) has not yet resolved the issues raised in your citizen petition received on October 21, 2013, and submitted on behalf of Covis Pharma Sàrl (Covis). Your petition requests that the Agency (1) require all sponsors of generic 0.0625 mg and 0.1875 mg strength digoxin tablets relying upon Lanoxin as the RLD to conduct and pass the same validation testing (dissolution and blend uniformity) that Petitioner conducted, and (2) deny any request from generic drug sponsors to waive the requirements for such testing.

FDA has been unable to reach a decision on your petition because it raises significant issues requiring extensive review and analysis by Agency officials. This interim response is provided in accordance with FDA regulations on citizen petitions (21 CFR 10.30(e)(2)). We will respond to your petition as soon as we have reached a decision on your request.

Sincerely,

Jane A. Axelrad

Associate Director for Policy

Center for Drug Evaluation and Research