

Food and Drug Administration 10903 New Hampshire Avenue Silver Spring, MD 20993

February 5, 2014

Mr. Reginald Burgess % Mr. Garrett Skelly, ESQ 160 Centennial Way Ste 21 Tustin, CA 92780

Re: Citizen Petition – Docket Number FDA-2013-P-0949

Dear Mr. Burgess:

This is an interim response to the petition dated August 6, 2013, filed by the Food and Drug Administration (FDA) on 8/9/2013. In the petition, you requested FDA to amend the clearance of the Otto Bock C-Leg, submitted under 510(k) premarket notification K991590, to include both prescription and over-the-counter use. Your petition further requests that all prosthetic devices regulated under Title 21 of the Code of Federal Regulations (CFR) 890.3420 and 890.3500 be labeled for over-the-counter use.

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

If you have any questions about this interim response, please contact Mr. Madhusoodana Nambiar of our Regulations Staff at (301) 796-5837.

Sincerely yours,

Nancy Stade

Deputy Director for Policy

Center for Devices and

Radiological Health