Date Posted: December 15, 2015
Recall Status: Open
Recall Number: Z-0394-2016
Recall Event ID: 72635
Premarket Notification 510(k) Number: K122953

Product Classification: Pump, Infusion, Insulin - Product Code LZG

Product: OmniPod®, Insulin Management System (OUS) Catalog Number: 14810 Product Usage: The OmniPod® Insulin Management System is intended for subcutaneous delivery of insulin at set and variable rates for the management of diabetes mellitus in persons requiring insulin and for the quantitative measurement of glucose in fresh whole capillary blood (in vitro) from the finger.

Code Information: Lot Codes: L41908, L41910, F41935
Recalling Firma/Manufacturer: Insulet Corporation 600 Technology Park Dr Ste 200 Billerica, Massachusetts 01821-4126

For Additional Information Contact: Same 978-600-7000
Manufacturer Reason for Recall: Pod's needle mechanism fails to deploy or there is a delay in the deployment of the needle mechanism.
FDA Determined Cause: CHANGE CONTROL (GMP - GOOD MANUFACTURING PRACTICE): Process Change Control

Action: Insulet issued on 11/2/15 an URGENT: Field Safety Notification via Email notification and Federal Express. Letter describes the problem that certain lots of the pod's needle failed to deploy or there is a delay in the deployment of the needle. Customers not responding to the email or Federal Express will receive additional mailing and/or follow up phone calls. Call Customer Care at 1-855-407-3729 if you have any questions regarding this Field Safety Notification.

Quantity in Commerce: 5,179.0 boxes
Distribution: Worldwide Distribution - US Nationwide and countries of Switzerland, Germany, and Israel.
Total Product Life Cycle: TPLC Device Report

1 For details about termination of a recall see Code of Federal Regulations (CFR) Title 21 §7.55
2 Per FDA policy, recall cause determinations are subject to modification up to the point of termination of the recall.