NP Thyroid® (Thyroid Tablets, USP)

Date: 05/22/2020

At Magellan Rx Management, we want to help you get the best possible care. We have created a site to share drug recall information.

Acella Pharmaceuticals, LLC has posted a lot recall of NP Thyroid® (Thyroid Tablets, USP) 30 mg, 60 mg, and 90 mg.

About this Recall:
Acella Pharmaceuticals, LLC is voluntarily recalling a total of 13 lots of 30 mg, 60 mg, and 90 mg NP Thyroid® (thyroid tablets, USP) to the consumer level. The products are being recalled because our testing has found these lots to be superpotent. The product may have up to 115% of the labeled amount of liothyronine (T3).

What this means to you:
Patients being treated for hypothyroidism (underactive thyroid), who receive superpotent NP Thyroid, may experience signs and symptoms of hyperthyroidism (overactive thyroid) which include, but are not limited to, weight loss, heat intolerance, fatigue, muscle weakness, high blood pressure, chest pain, rapid heart rate, or heart rhythm disturbances. Pregnant women who take superpotent NP Thyroid may also experience negative maternal and fetal outcomes including miscarriage and/or impairment of fetal development. The FDA is recommending patients who are currently taking NP Thyroid from a lot being recalled should not discontinue use without contacting their healthcare provider for further guidance and/or a replacement prescription. View the FDA Recall Notification at the link below for a listing of lots being recalled.

Consumers with questions about the recall can email Acella Pharmaceuticals at recall@acellapharma.com or contact Acella Customer Service at 1-800-541-4802, Monday through Thursday from 9:00 am to 5:00 pm ET and Friday from 9:00 am to 12:30 pm ET. Consumers should contact their physician or healthcare provider if they have experienced any problems that may be related to taking or using this drug product.

For more information regarding this FDA Recall Notification, please refer to the FDA website:


FDA contact information for reporting adverse events/quality complaints can be reached online at https://www.accessdata.fda.gov/scripts/medwatch/index.cfm?action=reporting.home

or by calling the FDA at 1-888-INFO-FDA (1-888-463-6332) and then select prompt #2.